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**Exceptional skull of huayqueriana (mammalia, litopterna, macraucheniidae)
from the late miocene of argentina: anatomy, systematics, and
paleobiological implications**

Forasiepi, Analia M ; MacPhee, R D E ; del Pino, Santiago Hernández ; Schmidt, Gabriela I ; Amson, Eli ; Grohé, Camille

Abstract: The Huayquerías Formation (Late Miocene, Huayquerian SALMA) is broadly exposed in westcentral Argentina (Mendoza). The target of several major paleontological expeditions in the first half of the 20th century, the Mendozan Huayquerías (“badlands”) have recently yielded a significant number of new fossil finds. In this contribution we describe a complete skull (IANIGLA-PV 29) and place it systematically as *Huayqueriana* cf. *H. cristata* (Rovereto, 1914) (Litopterna, Macraucheniidae). The specimen shares some nonexclusive features with *H. cristata* (similar size, rostral border of the orbit almost level with distal border of M3, convergence of maxillary bones at the level of the P3/P4 embrasure, flat snout, very protruding orbits, round outline of premaxillary area in palatal view, and small diastemata between I3/C and C/P1). Other differences (e.g., lack of sagittal crest) may or may not represent intraspecific variation. In addition to other features described here, endocast reconstruction utilizing computer tomography (CT) revealed the presence of a derived position of the orbitotemporal canal running below the rhinal fissure along the lateroventral aspect of the piriform lobe. CT scanning also established that the maxillary nerve (CN V2) leaves the skull through the sphenoorbital fissure, as in all other litopterns, a point previously contested for macraucheniids. The angle between the lateral semicircular canal and the plane of the base of the skull is about 26°, indicating that in life the head was oriented much as in modern horses. Depending on the variables used, estimates of the body mass of IANIGLA-PV 29 produced somewhat conflicting results. Our preferred body mass estimate is 250 kg, based on the centroid size of 36 3D cranial landmarks and accompanying low prediction error. The advanced degree of tooth wear in IANIGLA-PV 29 implies that the individual died well into old age. However, a count of cementum lines on the sectioned left M2 is consistent with an age at death of 10 or 11 years, younger than expected given its body mass. This suggests that the animal had a very abrasive diet. Phylogenetic analysis failed to resolve the position of IANIGLA-PV 29 satisfactorily, a result possibly influenced by intraspecific variation. There is no decisive evidence for the proposition that *Huayqueriana*, or any other litoptern, were foregut fermenters.

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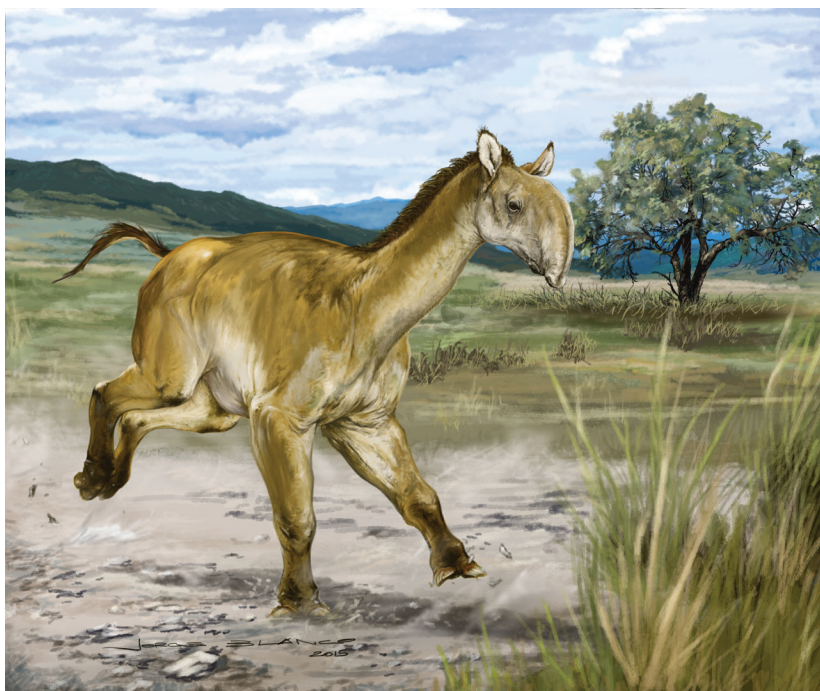
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(MAMMALIA, LITOPTERNA, MACRAUCHENIIDAE)
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ANATOMY, SYSTEMATICS, AND PALEOBIOLOGICAL
IMPLICATIONS

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BULLETIN OF THE AMERICAN MUSEUM OF NATURAL HISTORY

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ABSTRACT

The Huayquerías Formation (Late Miocene, Huayquerian SALMA) is broadly exposed in west-central Argentina (Mendoza). The target of several major paleontological expeditions in the first half of the 20th century, the Mendozaan Huayquerías (“badlands”) have recently yielded a significant number of new fossil finds. In this contribution we describe a complete skull (IANIGLA-PV 29) and place it systematically as *Huayqueriana* cf. *H. cristata* (Rovereto, 1914) (Litopterna, Macrauchenidae). The specimen shares some nonexclusive features with *H. cristata* (similar size, rostral border of the orbit almost level with distal border of M3, convergence of maxillary bones at the level of the P3/P4 embrasure, flat snout, very protruding orbits, round outline of premaxillary area in palatal view, and small diastemata between I3/C and C/P1). Other differences (e.g., lack of sagittal crest) may or may not represent intraspecific variation. In addition to other features described here, endocast reconstruction utilizing computer tomography (CT) revealed the presence of a derived position of the orbitotemporal canal running below the rhinal fissure along the lateroventral aspect of the piriform lobe. CT scanning also established that the maxillary nerve (CN V₂) leaves the skull through the sphenoorbital fissure, as in all other litopterns, a point previously contested for macrauchenids. The angle between the lateral semicircular canal and the plane of the base of the skull is about 26°, indicating that in life the head was oriented much as in modern horses. Depending on the variables used, estimates of the body mass of IANIGLA-PV 29 produced somewhat conflicting results. Our preferred body mass estimate is 250 kg, based on the centroid size of 36 3D cranial landmarks and accompanying low prediction error. The advanced degree of tooth wear in IANIGLA-PV 29 implies that the individual died well into old age. However, a count of cementum lines on the sectioned left M2 is consistent with an age at death of 10 or 11 years, younger than expected given its body mass. This suggests that the animal had a very abrasive diet. Phylogenetic analysis failed to resolve the position of IANIGLA-PV 29 satisfactorily, a result possibly influenced by intraspecific variation. There is no decisive evidence for the proposition that *Huayqueriana*, or any other litoptern, were foregut fermenters.

INTRODUCTION

By any measure, the morphological diversification of South American native ungulates (SANUs) during the Cenozoic was extraordinary. In size alone, they ranged from notoungulates no larger than rabbits (e.g., *Pachyrukhos*, ~2 kg; Cassini et al., 2012a) to rhino-sized astrapotheres (e.g., *Parastrapotherium*, ~2000 kg; Kramarz and Bond, 2008). In terms of taxonomic diversity, Notoungulata was the most successful of the SANU orders, followed by Litopterna with about a hundred species discriminated at present (Cifelli, 1983, 1993; Cifelli and Soria, 1983; Bond, 1986, 1999; Bond et al., 1995, 2001; Cifelli and Guerrero, 1997; Soria, 1981, 2001; Scherer et al., 2009; Schmidt and Ferrero, 2014). Nevertheless, despite their numerous morphological specializations, according to collagen proteomic studies these two SANUs not only appear to have shared

a common ancestry but also are most closely related to Perissodactyla among major extant placental taxa (Welker et al., 2015; Buckley, 2015). The relationships of other major SANU clades to one another or to other placentals have not been resolved.

Several families have been classically incorporated within Litopterna. The central families arose at the end of the Oligocene and extensively radiated during the Neogene: Adianthidae (Late Oligocene–Early Miocene), Protheroheriidae (Late Oligocene–Late Pleistocene), and Macrauchenidae (Late Oligocene–Late Pleistocene/Early Holocene), the latter represented by Cramaucheniinae (including Theosodontinae) and Macraucheniiinae (Cifelli, 1983; Bond, 1986, 1999; Soria, 1986, 2001; Schmidt and Ferrero, 2014). Various Paleogene groups (Amilnedwardsiidae, Anisolambdidae, Indalecidae, Notonychopidae,

TABLE 1
Checklist of the family Macraucheniidae

<i>Cullinia levis</i>	Cabrera and Kraglievich, 1931	Late Miocene
<i>Huayqueriana cristata</i>	(Rovereto, 1914) ¹	Late Miocene
<i>Macrauchenia patachonica</i>	Owen, 1838	Quaternary
<i>Macrauchenopsis ensenadensis</i>	(Ameghino, 1888) ²	Quaternary
<i>Oxyodontherium zeballosi</i>	Ameghino, 1883a ³	Late Miocene
<i>Paranauchenia hystata</i>	(Cabrera and Kraglievich, 1931)	Late Miocene
<i>Paranauchenia denticulata</i>	(Ameghino, 1891a)	Late Miocene
<i>Promacrauchenia antiquua</i>	(Ameghino, 1887a) ⁴	Early Pliocene
<i>Promacrauchenia calchaquiorum</i>	Rovereto, 1914	?Late Miocene / ?Early Pliocene
<i>Promacrauchenia chapadmalense</i>	Ameghino, 1908	Early Pliocene
<i>Promacrauchenia kraglievichi</i>	Parodi, 1931	Early Pliocene
<i>Promacrauchenia (Pseudomacrauchenia) yepesi</i>	Kraglievich, 1930	Quaternary
<i>Scalabrinittherium bravardi</i>	Ameghino, 1883b ⁵	Late Miocene
<i>Scalabrinittherium rusconi</i>	Parodi, 1931 ⁶	?Late Miocene / ?Pliocene
<i>Windhausenia delacroixi</i>	Kraglievich, 1930	Late Pliocene
<i>Xenorhinotherium bahiense</i>	Cartelle and Lessa, 1988	Quaternary

¹ Includes *Macrauchenidia latidens* Cabrera, 1939.
² Not certainly valid, may be part of *Windhausenia* (e.g., Marshall et al., 1984).
³ Includes *Mesorhinus* Ameghino, 1885.
⁴ See text fn. 1.
⁵ Includes *Scalabrinia* Lydekker, 1894.
⁶ Not certainly valid.

Protolipternidae, and Sparnotheriodontidae) are also frequently cited as true litopterns, but whether they actually belong in the same monophyletic group as the later taxa is debatable (Cifelli, 1993; Billet et al., 2015).
Macraucheniinae currently includes 16 species (table 1), some of which are based on isolated postcranial material and are of uncertain validity (e.g., Parodi, 1931; Kraglievich, 1930; Paula Couto, 1945). At least from the standpoint of taxonomic richness, peak diversity for this group was achieved in the Late Miocene, with some localities containing as many as five coeval species (e.g., Ituzaingó Formation; Schmidt and Cerdeño, 2013). Species richness on this scale may be regarded as due either to particularly intense selection and environmental exploitation by contemporaneous taxa, or to

taxonomic inflation consequent on limited or noncomparable hypodigms.
The macraucheniine fossil record is comparatively good, particularly for the Pleistocene (Fernández de Álvarez, 1940). *Macrauchenia patachonica*, the first remains of which were recovered by Charles Darwin on January 9, 1834, at Puerto San Julian in Patagonia, was also the last member of the family to disappear, persisting in the Pampean region of Argentina into the Early Holocene (8390 ± 140 ¹⁴C yr BP; Tonni, 1990; Bond, 1999).
Macraucheniines were long-necked, horse-sized herbivores, possibly cursorial with three-digit cheiridia. In this subfamily the molars are encased in cementum and mesodont, with deeper enamel fossettes than in other macraucheniids (e.g., *Cramauchenia*, *Theosodon*) (Bond, 1999).

This tooth morphology probably evolved to cope with the hard parts of xerophytic plants (e.g., thick foliar cuticles and spines, fruits encased in hard shells or rinds to combat desiccation; Villagra et al., 2011) typical of the dry, open environments that became the rule in much of southern South America as a consequence of Late Miocene climate change. Decreasing global temperatures, together with greater seasonality, aridity, and the spread of C4 grasslands (Cerling et al., 1997; Zachos et al., 2001) were correlated with an increase in the diversity and abundance of mammals with hypsodont teeth (Jernvall and Fortelius, 2002; Eronen et al., 2010). Hypsodonty is particularly marked in the South American mammal record from the mid-Paleogene onward (Madden, 2015). Euhypsodonty (= elodonty; ever-growing, open-rooted teeth) is very widespread in Late Miocene herbivores (e.g., sloths, armadillos, glyptodonts, notoungulates, and many caviomorph rodents), although litopterns, unlike notoungulates, never developed this adaptation. On the basis of measurements of the temporal fossa in *Theosodon*, Madden (2015) has hypothesized that at least some macraucheniiids may have been foregut fermenters, like artiodactyl ruminants. However, the evidence is not persuasive (see Digestive Physiology).

One of the most characteristic features of the macraucheniiine skull is the repositioning of the nasal aperture. Instead of being situated at the front of the face, below the orbits and immediately above the incisors, the aperture is situated in a centrodorsal position, between the eyes, near the morphological summit of the skull. Although it has long been assumed that this highly derived positioning implies the existence of a trunk or similar appendage (Burmeister, 1864; Scott, 1937; Rusconi, 1957; Soria, 1981; Bond, 1999), detailed comparative anatomical evaluations in support of this inference are lacking.

In this contribution, we provide a detailed comparative description of an exceptionally well-preserved skull of *Huayqueriana* cf. *H. cristata* (IANIGLA-PV 29) from the Late Miocene Huayquerías Formation, Mendoza, Argentina, utilizing

both traditional methods of description as well as micro-CT scanning. We additionally evaluate species taxonomy, phylogenetic relationships, and paleobiological parameters such as body mass, longevity, and digestive physiology.

GEOGRAPHICAL AND GEOLOGICAL CONTEXTS

The huayquerías, or badlands, of central Mendoza (fig. 1A) have yielded fossils of a wide range of mammalian taxa generally regarded as Late Miocene–Pliocene in age (Marshall et al., 1986; Yrigoyen, 1994). Paleontological studies in the area were initiated in the first half of the 20th century and have continued on an irregular basis since then (De Carles, 1911; Rovereto, 1914; Frenguelli, 1930; Rusconi, 1939; Marshall et al., 1986; Forasiepi et al., 2014, 2015a).

The Huayquerías del Este (Eastern Badlands) present three outcropping formations. These include a Late Miocene unit, (1) the Huayquerías Formation, and two Pliocene units, (2) the Tunuyán and (3) the Bajada Grande formations (Dessanti, 1946; Marshall et al., 1986; Yrigoyen, 1993, 1994). The Huayquerías del Este are dominated by a large, extensively faulted anticline, in the central portion of which the eponymous Huayquerías Formation outcrops (to avoid possible confusion between geographical and formational names, the Huayquerías Formation will hereafter be abbreviated to HF). The HF is ~200 m thick (Marshall et al., 1986) and is composed of massive beds of brownish-grey sandstones with sporadic intercalations of red mudstones (fig. 1B; Dessanti, 1946; Marshall et al., 1986), deposited in an alluvial environment with ephemeral rivers and muddy floodplains (Garrido et al., in prep.; Forasiepi et al., 2015a).

A fourth unit, the Tobas Angostura Formation, does not outcrop in the study area but does so farther north. Yrigoyen (1993) inferred that this unit, which has an estimated age of 10.5–9.5 Ma on the basis of K/Ar dating, must immediately underlie the HF. If that inference is correct, the base of the HF cannot be much less than the

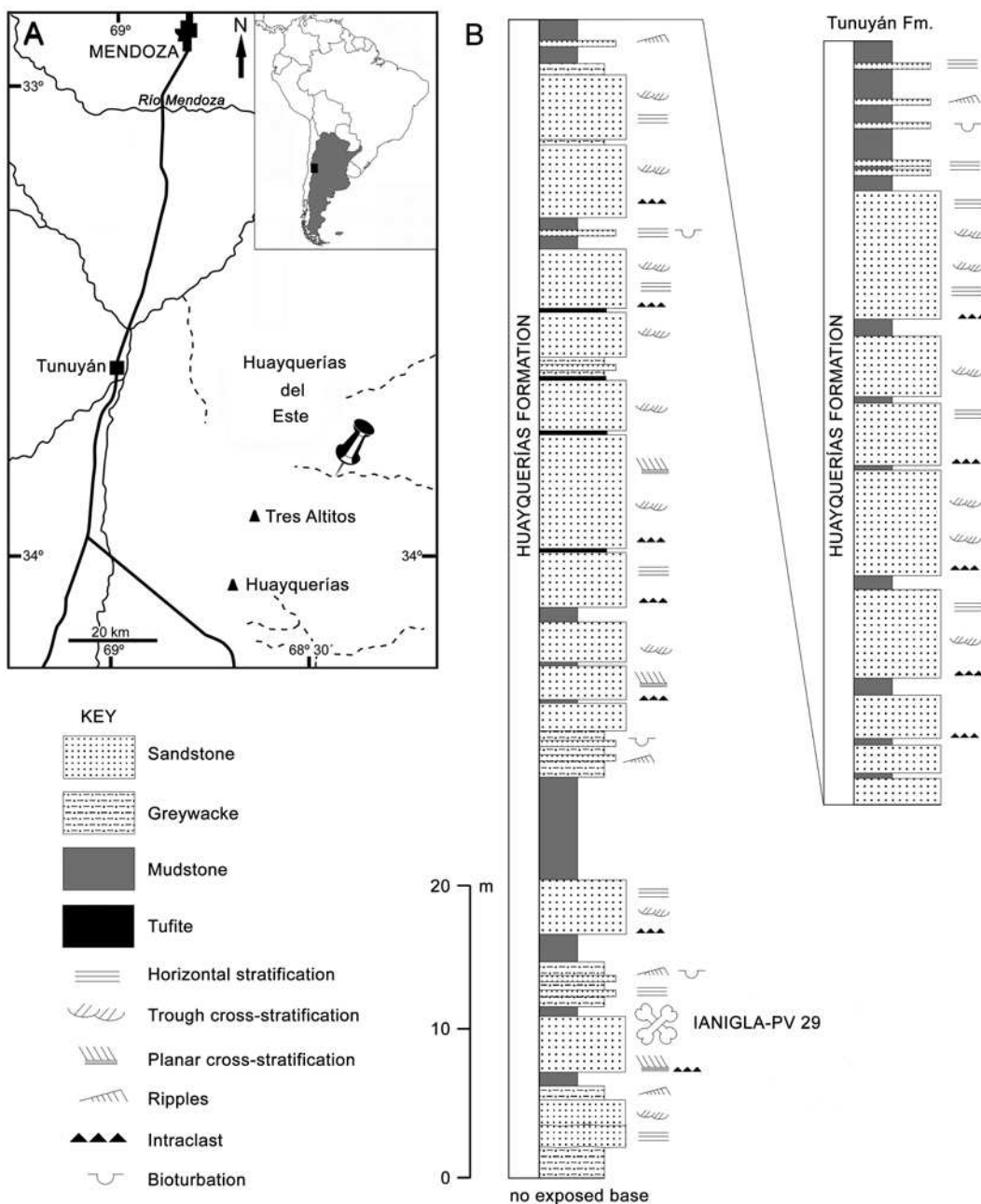


FIG. 1. Geographical and stratigraphic context: **A**, location of Río Seco de la Última Aguada, north-central Mendoza, Argentina; **B**, integrated stratigraphic column, Río Seco de la Última Aguada, Huayquerías del Este. *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29 was recovered from lower levels of Huayquerías Formation (Late Miocene; Huayquerian) exposed along river gorge.

indicated range. The top of the HF was estimated to be approximately 5.8 Ma by Yrigoyen (1994) on the basis of K/Ar dates reported by Marshall et al. (1986). However, because of uncertainties concerning the stratigraphic extent and definition of the overlying Tunuyán Formation (Garrido et al., in prep.), this estimate has to be treated with caution.

The Huayquerian South American Land Mammal Age (SALMA) was originally defined on the basis of a small series of mammal taxa from HF as then understood (Kraglievich, 1934; Simpson, 1940). The original faunal list included *Megatheriops rectidens* (Megatheriidae), *Proscelidodon gracillimus* (Mylodontidae), *Hemihegetotherium achataleptum* (Hegetotheriinae), *Neobrachytherium* sp. (Protherotheriidae), *Huayqueriana cristata* (Macraucheniiidae), *Cyonasua pascuali* (Procyonidae), and *Lagostomus pretrichodactyla* (Chinchillidae) (Rovereto, 1914; Rusconi, 1939; Kraglievich and Olazábal, 1959; Linares, 1981; Marshall et al., 1983; Pascual and de la Fuente, 1993; Soria, 2001). It is important to note that Huayquerian age assignments have frequently been made for Neotropical mammal associations recovered from a wide variety of other contexts throughout South America, such as Urumaco in Venezuela (Linares, 2004) and Acre in Brazil (Cozzuol, 2006; Ribeiro et al., 2013). This also applies to the correlation of presumed age-equivalent stratigraphic units, such as the Cerro Azul and Epecuén Formations in the Pampean area (Goin et al., 2000; Montalvo et al., 2008; Verzi et al., 2011), the Chiquimil or Andahuala formations in northwestern Argentina (e.g., Reguero and Candela, 2011; see also Esteban et al., 2014, for a discussion on the topic and references herein cited). However, it should be noted that the evidence for some of these faunal and temporal correlations with the Huayquerian SALMA as presently understood is notably slim.

Since 2013, new field work has been conducted in the HF in order to more fully characterize its actual faunal content, and thus lay a basis for a critical reassessment of the definition of the Huayquerian SALMA. Results to date

include the first fossil footprints of a SANU (aff. *Macrauchenichnus* isp.) to be identified in this formation, as well as a notable expansion of the faunal list from the original seven taxa to 20+ (including additional mammal, reptile and anuran taxa currently under study; Forasiepi et al., 2014, 2015a).

Among our fossil discoveries is the subject of this paper, IANIGLA-PV 29, a beautifully preserved skull here referred to *Huayqueriana* cf. *H. cristata*. This specimen was collected from the HF sediments forming the floor of the Río Seco de la Última Aguada (68°27'31.6"W, 33°54'14.5"S), on the southeast side of the Huayquerías del Este (fig. 1B). According to De Carles (1911), the holotype of *Huayqueriana cristata* (MACN-PV 8463; fig. 2) was collected in the Huayquerías de Biluco, in the vicinity of Agua de la Concha at the foot of Cerro Torrecitas. This raises an interesting biostratigraphic issue, because the Tunuyán (nominally Pliocene) and the Huayquerías (nominally Late Miocene) formations both outcrop in this area (Yrigoyen, 1994). In addition, the sediment associated with MACN-PV 8463, a tuffaceous sandstone with fine-grained, immature epiclasts formed by quartz, hornblende, and other inclusions that react strongly to HCl, is more reminiscent of the Tunuyán Formation than the HF (A.C. Garrido, personal commun., 2015).

MATERIAL AND METHODS

Descriptions of IANIGLA-PV 29 in the following pages are grouped by region: cranium (face, orbit, vault, zygoma, basicranium), cerebral endocast, and upper dentition. Anatomical terminology generally conforms to, or is adapted from, nomina recognized in *Terminologia Anatomica* (e.g., Schaller, 1992); for more specialized terms, we utilized MacPhee (1981, 2014) for the caudal cranium, Soria (2001) for the dentition, and Macrini et al. (2007a, 2007b) for the cerebral endocast. The description of the inner ear assumes that the lateral semicircular canal is oriented horizontally and centrally within the surrounding

array of anatomical structures (Graf and Klam, 2006; Ekdale, 2013). Comparisons to other macraucheniids are based on direct study of specimens as well as descriptions and illustrations in the literature (appendix 1). Museum collection numbers are provided in the text when more than one individual is known for a single taxon.

In view of recent proteomic evidence for the close relationship of litopterns and notoungulates to Perissodactyla (Welker et al., 2015; Buckley, 2015), we make specific comparisons to living horses, tapirs, and rhinos where warranted. However, our primary purpose here is to investigate the relationships of *Huayqueriana* within the context of Litopterna, not to test the likelihood of Panperissodactyla (Welker et al., 2015) from the standpoint of cranial morphology.

Segmental data on *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29 and *Tetramerorhinus lucarius* AMNH 9245 were collected with the GE Phoenix vtomex s240 μ CT scanner housed at AMNH. The skull of *Huayqueriana* was scanned at 180 kV, 220 mA, with a voxel size of 124.06 μ m. The scan resulted in a final stack of 3585 slices of 1619 \times 1166 pxl. The program 3DSlicer (open source <http://www.slicer.org/>; Fedorov et al., 2012) was used for 3D reconstruction and segmentation of the skull, endocast, and associated structures. Mimics 18.0 (Materialise NV, Leuven, Belgium) was used for 3D reconstruction of the petrosal and VGStudio MAX (Volume Graphics GmbH, Heidelberg, Germany) for the osseous inner ear. Linear and volumetric measurements of the endocast were calculated with 3D Slicer following standard dimensions proposed by Macrini et al. (2007b); angular measurements were calculated with the program ImageJ; linear measurements of the bony labyrinth were calculated with Mimics following the protocol of Schmelzle et al. (2007) for the dimensions of the semicircular canals. Height and width of the cochlea were measured in profile.

The left M2 of IANIGLA-PV 29 was extracted, embedded in Araldite epoxy resin, and prepared for histological study by cutting (1) a coronal section, at the level of the interroot pad where the cementum is usually the thickest, and (2) a

paraocclusal section, in the distal root, apical to the crown. Cut surfaces were polished with Al₂O₃ sandpapers with successively smaller grit sizes (320 to 25,000). The polished surfaces were lightly coated with glycerine, and observed with a stereoscopic microscope (Leica MZ 16⁺) under natural light. Images were taken with a Leica DFC 420 C⁺ digital camera, and their contrast was enhanced with Adobe Photoshop[®]. For comparative purposes, the M1 of *Theosodon garrettorum* (PIMUZ A/V 4662) was subjected to the same procedure.

Phylogenetic relationships were explored via a cladistic analysis employing TNT (Goloboff et al., 2008). The data matrix is a modified version of that published by Schmidt and Ferrero (2014).

The body mass of IANIGLA-PV 29 was estimated using a variety of allometric equations in the literature that are in turn based on dental and cranial traits of extant euungulates (Artiodactyla and Perissodactyla) (see Body Size). Of general interest is our finding that body mass estimators based on cranial measurements were more plausible than ones based on teeth (cf. Cassini et al., 2012b).

ABBREVIATIONS

INSTITUTIONAL: AMNH, American Museum of Natural History, New York. IANIGLA-PV, Instituto Argentino de Nivología, Glaciología y Ciencias Ambientales, Vertebrate Paleontology, Mendoza, Argentina. MACN, Museo Argentino de Ciencias Naturales “Bernardino Rivadavia” (A, Ameghino; PV, Vertebrate Paleontology; SC Santa Cruz collections), Buenos Aires, Argentina. MHIN-UNSL-GEO-V, Museo de Historia Natural, Departamento de Geología of the Universidad Nacional de San Luis, Paleontological Collection, San Luis, Argentina. MLP, Museo de La Plata, Buenos Aires, Argentina. MPEF-PV, Museo Paleontológico “Egidio Feruglio,” Trelew, Chubut, Argentina. NMB, Naturhistorisches Museum Basel, Switzerland. PIMUZ, Paläontologisches Institut und Museum, Universität Zürich, Switzerland; ZMUZH, Zoologisches Museum der Universität Zürich, Switzerland.

ANATOMICAL: Abbreviations used in individual figures, most of which are set in lower-case, are indicted in captions. In the main text, we use additional conventions. Upper teeth are capitalized while mandibular teeth are in lower-case, as follows: C/c, canine, I/i incisor; M/m, molar; and P/p, premolar; D/d, deciduous dentition. For the petrosal, we capitalize the following abbreviations: IAM, internal acoustic meatus; CTPP, caudal tympanic process of the petrosal; FAI, inferior acoustic foramen; FAS, superior acoustic foramen, RTPP, rostral tympanic process of petrosal; SIPS, sulcus for the inferior petrosal sinus. For the inner ear: ASC, anterior semicircular canal; LSC, lateral semicircular canal; PSC, posterior semicircular canal; L, left; R, right.

OTHER ABBREVIATIONS: CI, consistency index; RI, retention index; TBR, tree bisection reconnection.

SYSTEMATIC PALEONTOLOGY

ORDER Litopterna Ameghino, 1889

FAMILY Macrauchiidae Gervais, 1855

SUBFAMILY Macrauchiinae Gervais, 1855

Huayqueriana Kraglievich, 1934

Huayqueriana Kraglievich, 1934: 117 (as subgenus of *Promacrauchenia*).

Huayqueria Riggs and Patterson, 1939: 157 (*lapsus calami*).

Macrauchenidia Cabrera, 1939: 167

TYPE SPECIES: *Promacrauchenia? cristata* Rovereto, 1914.

Huayqueriana cristata (Rovereto, 1914)

Promacrauchenia? cristata Rovereto, 1914: 212–213; pl. 29, figs. 4, 4a.

Promacrauchenia (Huayqueriana) cristata Kraglievich, 1934: 117.

Huayqueria [sic] *cristata* Riggs and Patterson, 1939: 157.

Macrauchenidia latidens Cabrera, 1939: 167: 33–34, fig. 24; Pascual, 1966, pl. 71, figs. D and E.

Huayqueriana cristata Soria, 1986: 158.

HOLOTYPE: MACN-PV 8463 (fig. 2), fragmentary skull and associated dentaries.

Huayqueriana cf. *H. cristata* (Rovereto, 1914)

REFERRED SPECIMEN: IANIGLA-PV 29, cranium with LP1–3, distal half of LP4, RP2–4, and R/LM2–3 (figs. 3, 4, 10–22, 24–25, 27–28; tables 2–4).

GEOGRAPHIC AND STRATIGRAPHIC DISTRIBUTION: Huayquerías del Este, Río Seco de la Última Aguada (68°27'31.6"W, 33°54'14.5"S), San Carlos Department, Mendoza Province, Argentina (fig. 1A). Huayquerías Formation (fig. 1B), Huayquerian SALMA (currently correlated with Tortonian-Messinian stages of Late Miocene; Marshall et al., 1986; Yrigoyen, 1994).

MORPHOLOGY OF IANIGLA-PV 29, *HUAYQUERIANA* CF. *H. CRISTATA*

CRANIUM

The skull is long, narrow, and low (fig. 3A). In overall size the specimen is similar to the holotype of *Huayqueriana cristata* (fig. 2), *Promacrauchenia antiquua*,¹ *Promacrauchenia calchaquiorum*, and some specimens of *Scalabrinittherium bravardi* (e.g., MACN-A 1270); overall it is slightly larger than *Oxydontherium zeballosi*, but smaller than *Windhausenienia delacroixi*, *Macrauchenia patachonica*, and *Xenorhinotherium bahiense* (table 2; fig. 4). Most cranial elements are completely fused, with sutures obliterated; muscular scars are deep and teeth are heavily worn. Several teeth were lost during life, as indicated by bony infill within alveoli. These features strongly indicate that IANIGLA-PV 29 was a senile individual when it died (see Longevity). Although tooth wear is a proximate indicator of senescence (Skogland, 1988; Loe et al., 2003), wear rate will depend on dietary regime.

¹ Ameghino (1887a) described *Macrauchenia antiqua* as a new species from the Montehermosan of Buenos Aires. Since the trivial name is not spelled “antigua,” the modern Spanish spelling of the cognate Latin fem. sing. adj. “antiquua,” we infer that Ameghino’s intention was clear but that he inadvertently substituted the former for the latter. Under the provisions of ICZN art. 33.2, we emend this name to *Promacrauchenia* (= *Macrauchenia*) *antiquua*.

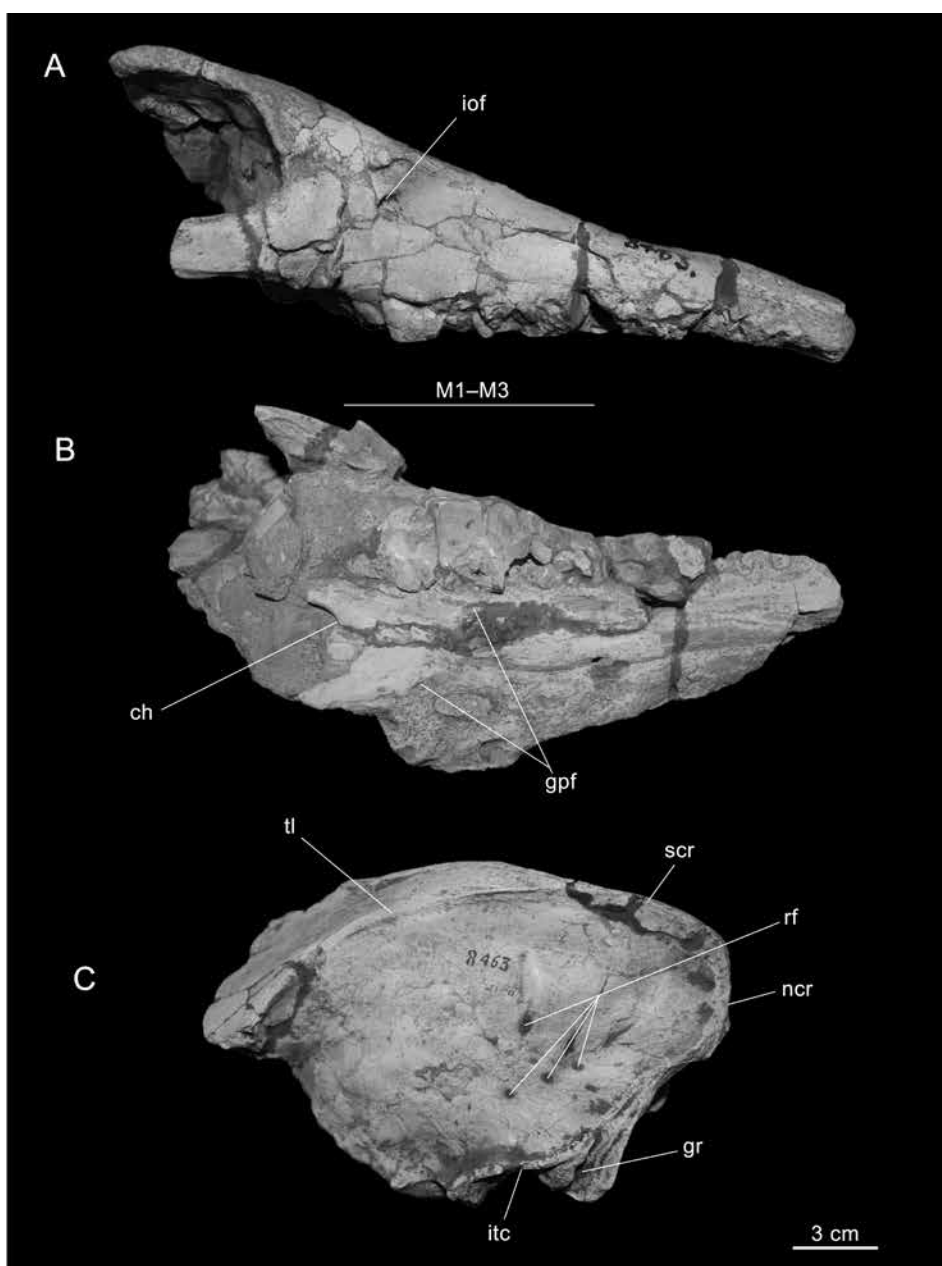


FIG. 2. Holotype of *Huayqueriana cristata* MACN-PV 8463 referred to Huayquerías Formation (Late Miocene, Huayquerian SALMA). **A**, right lateral and **B**, ventral views of snout; **C**, portion of braincase in left lateral view. Abbreviations: **ch**, choana; **gpf**, greater (= major) palatine foramen; **gr**, occipital groove; **iof**, infraorbital foramen; **itc**, infratemporal crest; **M**, molar; **ncr**, nuchal crest; **rf**, foramina for rami temporales; **scr**, sagittal crest; **tl**, temporal line.

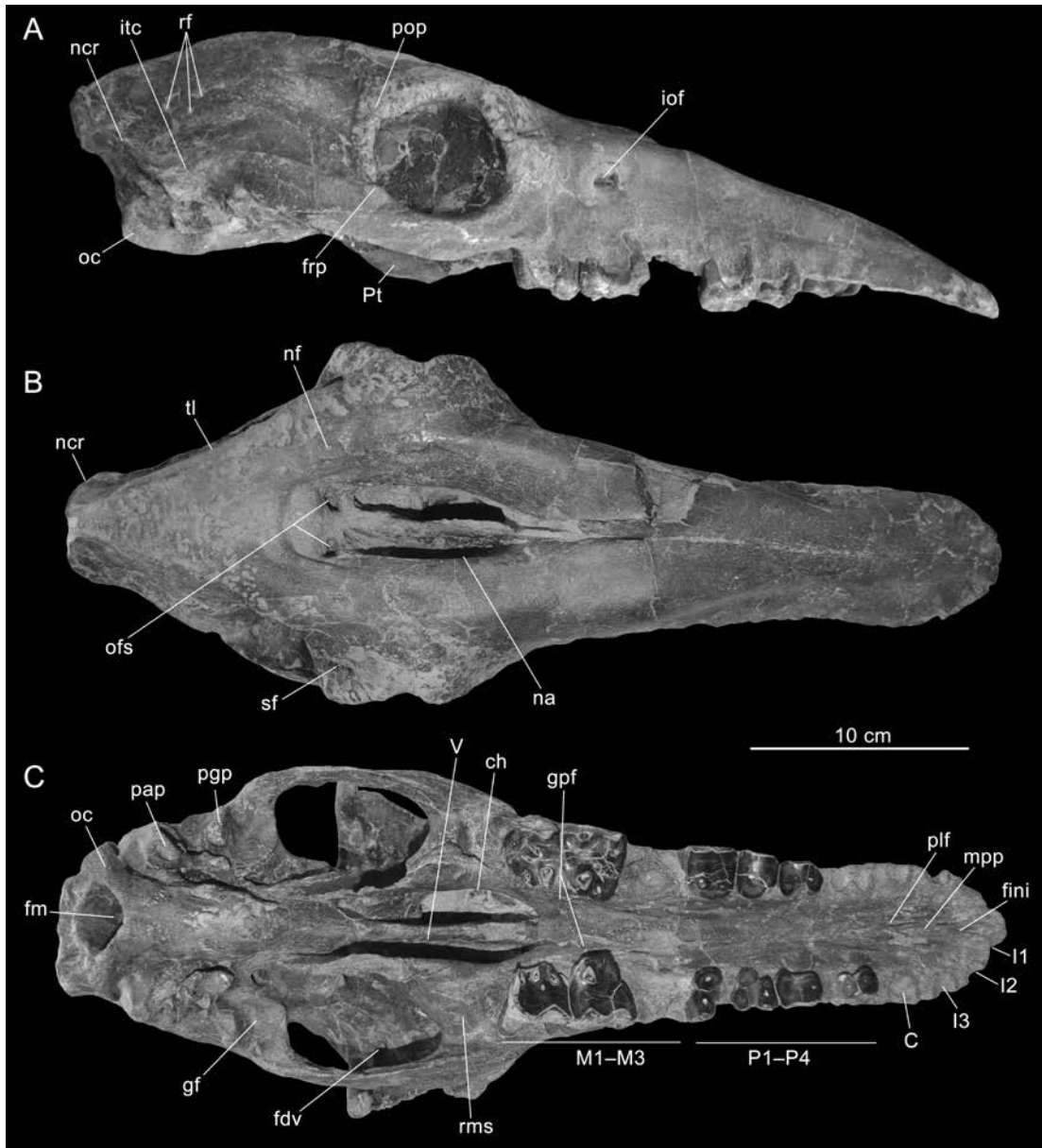


FIG. 3. Skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29 from Huayquerías Formation (Late Miocene, Huayquerian SALMA) in **A**, right lateral, **B**, dorsal, and **C**, ventral views. Abbreviations: **C**, canine (alveolus); **ch**, choana; **fdv**, foramen for frontal diploic vein; **fini**, foramen interincisivum; **fm**, foramen magnum; **frp**, frontal process of squamosal; **gf**, glenoid fossa; **gpf**, greater (= major) palatine foramen; **I**, incisor (alveolus); **iof**, infraorbital foramen; **itc**, infratemporal crest; **M**, molar; **mpp**, medial palatine process of premaxilla; **na**, nasal aperture; **ncr**, nuchal crest; **nf**, nutrient foramina; **oc**, occipital condyle; **ofs**, openings of frontal sinus; **P**, premolar; **pap**, paracondylar process of exoccipital; **pgp**, postglenoid process; **plf**, palatine fissure; **pop**, postorbital process of frontal; **Pt**, pterygoid; **rf**, foramina for rami temporales; **rms**, retromolar space; **sf**, supraorbital foramen; **tl**, temporal line; **V**, vomer.

Taphonomically, some elements of the cranium were broken or lost during burial, and the skull has suffered slight dorsolateral compression.

FACE

In lateral view, the narrow snout gradually increases in height from front to back (fig. 3A), much as in *Pr. antiquua* and *Pr. calchaquiorum*. The bone comprising the rostralmost part of the snout is very slightly ventrally curved, a feature also seen in *Pr. antiquua* (fig. 5B) and possibly the holotype of *H. cristata* (fig. 2A). Curvature is slightly more pronounced in *H. cristata* (MLP 41-IV-29-4; 5A), *O. zeballosi* (MACN-PV 13671; MHIN-UNSL-GEO-V 465a, fig. 8A), and *Pr. calchaquiorum* (fig. 5C), whereas in *S. bravardi* (MACN-A 1270), *M. patachonica*, and *X. bahiense* the equivalent area is planar.

The premaxilla/maxilla sutures are obliterated, and maxillae are partially fused midsagittally (fig. 3B). The roughly circular infraorbital foramen is large (~7 mm maximum diameter, compared to foramen ovale, 3.5 mm). This implies, although does not prove, that the parts of the face supplied by local neurosensory (CN V₂) and vascular bundles (infraorbital artery and vein) passing through the infraorbital foramen may have been of significant size, such as external nasal structures. The foramen opens dorsal to the transverse level of the distal part of M2 (fig. 3A), as in *H. cristata* (fig. 2A), *Pr. antiquua* (fig. 5B), *Pr. calchaquiorum* (fig. 5C), *Paranauchenia denticulata*, and *X. bahiense* (Cartelle and Lessa, 1988; Schmidt and Ferrero, 2014). The infraorbital foramen is positioned slightly more rostrally in *O. zeballosi* (MACN-PV 13671) where it opens above the M1/M2 embrasure; in MACN-PV 8903 (referred to *S. bravardi*), it is situated at the level of the mesial part of M2. By contrast, it is slightly more caudal in *M. patachonica* (right side of the skull MACN-PV 2; MLP 12-1424 and 12-1425), where it opens at the level of the M3 (left side of the skull MACN-PV 2 exhibits two infraorbital foramina, one at the level of the M2 and the other at the level of the M3). In IANIGLA-PV 29, the maxillary foramen (caudal

opening of the infraorbital canal) is of large size and compressed against the zygomatic arch.

The peculiar series of small sulci that issue from the infraorbital foramina, strongly marking both sides of the snout in specimens of *M. patachonica* (e.g., Burmeister, 1864), are also seen in *Pr. antiquua* MACN-PV 7986 (one only on each side), but are absent in *Pr. calchaquiorum*, MACN-PV 5528, and IANIGLA-PV 29. The face of *Scalabrinitherium* is missing and thus cannot be evaluated. The peculiarity of these sulci arises from the fact that, in *M. patachonica*, some of them actually overlap or cut through one another. They also seem to end blindly, with no reentry foramina within the maxilla. This suggests that their contents (presumably cutaneous nerves and associated blood vessels) were directed outward, toward large soft tissue targets in the face and snout.

NASAL APPARATUS

The outstanding feature of IANIGLA-PV 29 in dorsal view (fig. 3B) is the enormous nasal aperture, framed by the hypertrophied maxillae laterally and the diminutive nasals caudally. The aperture is situated near the center of the cranial roof, as in other macraucheniines (figs. 6, 8). The sloping margins of the aperture open directly into the relatively large choanae except caudally, where there is a low bony shelf flanked by two symmetrically positioned openings, as in *Pr. calchaquiorum* MACN-PV 5528 (fig. 6C) and *S. bravardi* MACN-PV 13082 (fig. 9B). In IANIGLA-PV 29 these openings represent aditus into the paranasal frontal sinuses (fig. 11) (see Pneumatization). In the other fossils we assume conditions were similar, although we cannot confirm this because the equivalent area is either damaged or not fully prepared.

Dorsally, the caudal border of the nasal aperture intersects a line joining the caudal borders of the orbits, as in other macraucheniines (Soria, 1986). In dorsal view, in general outline the nasal aperture is roughly teardrop shaped, widest between the orbits and gradually tapering rostrally to a point at which the two maxillary bones converge at the level of the P3/P4 embrasure (fig.

TABLE 2
Huayqueriana cristata (holotype, MACN-PV 8463) and *Huayqueriana* cf. *H. cristata* (IANIGLA-PV 29):
Cranial and dental measurements (in mm)^{1, 2, 3}

Skull	MACN-PV 8463	IANIGLA-PV 29
(1) Skull, total length (TSL)		435
(2) Skull rostrum, length from tip of premaxilla to posterior border of orbit		300
(3) Skull vault, length		150
(4) Skull, width at level of zygomatic arches		136
(5) Face, depth below ventral rim of orbit (SD)		112.04
(6) Zygomatic arch, maximum height at level of orbit		18.15
(7) Orbit, length		50.93
(8) Orbit, height		57.34
(9) Frontal, width at level of postorbital processes		153.9
(10) Nasal aperture, maximum width		30.13
(11) Nasal aperture, maximum length		145.01
(12) Muzzle, maximum width (MZW)		62.6
(13) Palate, length, lingual border incisor alveoli to distal border M3		212.05
(14) Palate, length, lingual border incisor alveoli to choana		190.15
(15) Posterior skull, length (PSL/SC)		169.65
(16) Palate, width at level of C		40
(17) Palate, width at level of P1		35.9
(18) Palate, width at level of P3		31.4
(19) Palate, width at level of M1		33.1
(20) Palate, width at level of M2 (PAW)		30.4
(21) Palate, width at level of M3	32	37
(22) Occiput, height (OCH)		83.34
(23) Supraoccipital, height, nuchal crest to foramen magnum		57.8
(24) Basicranium, length (BCL) ⁴		135.25 (*)
(25) Basicranium, width at the level of EAM	95 (*)	99.73
(26) Foramen magnum, height x width		24.6x26.3
Dentition		
(27) I1–M3, length (alveolus of I1)		215 (*)
(28) P1–M3, length		163
(29) P1–P4, length		86.7
(30) M1–M3, length	75(*)	82.2 (R) 79.1 (L)
P1 L		18.4 (L)
P1 W		11.5 (L)
P2 L		21.6 (R)
P2 W		16.7 (R)
P3 L		21.2 (R)
P3 W		21.1 (R)
P4 L		22.3 (R)
P4 W		22.6 (R)
M2 L (SUML)	30.2	31.4 (L)/30.9 (R)
M2 W (SUMW)	27.4	29.7 (L)/28.9 (R)
M3 L	25 (*)	26.1 (L)/26.3 (R)
M3 W	26.6	26.2 (L)/26 (R)

¹ Asterisk (*) indicates measurement is approximate.

² The numbers between parentheses relate to figure 4.

³ Abbreviations in parentheses relate to body size estimators in table 4.

⁴ From the base of the foramen magnum to the posterior border of choana. EAM, external acoustic meatus.

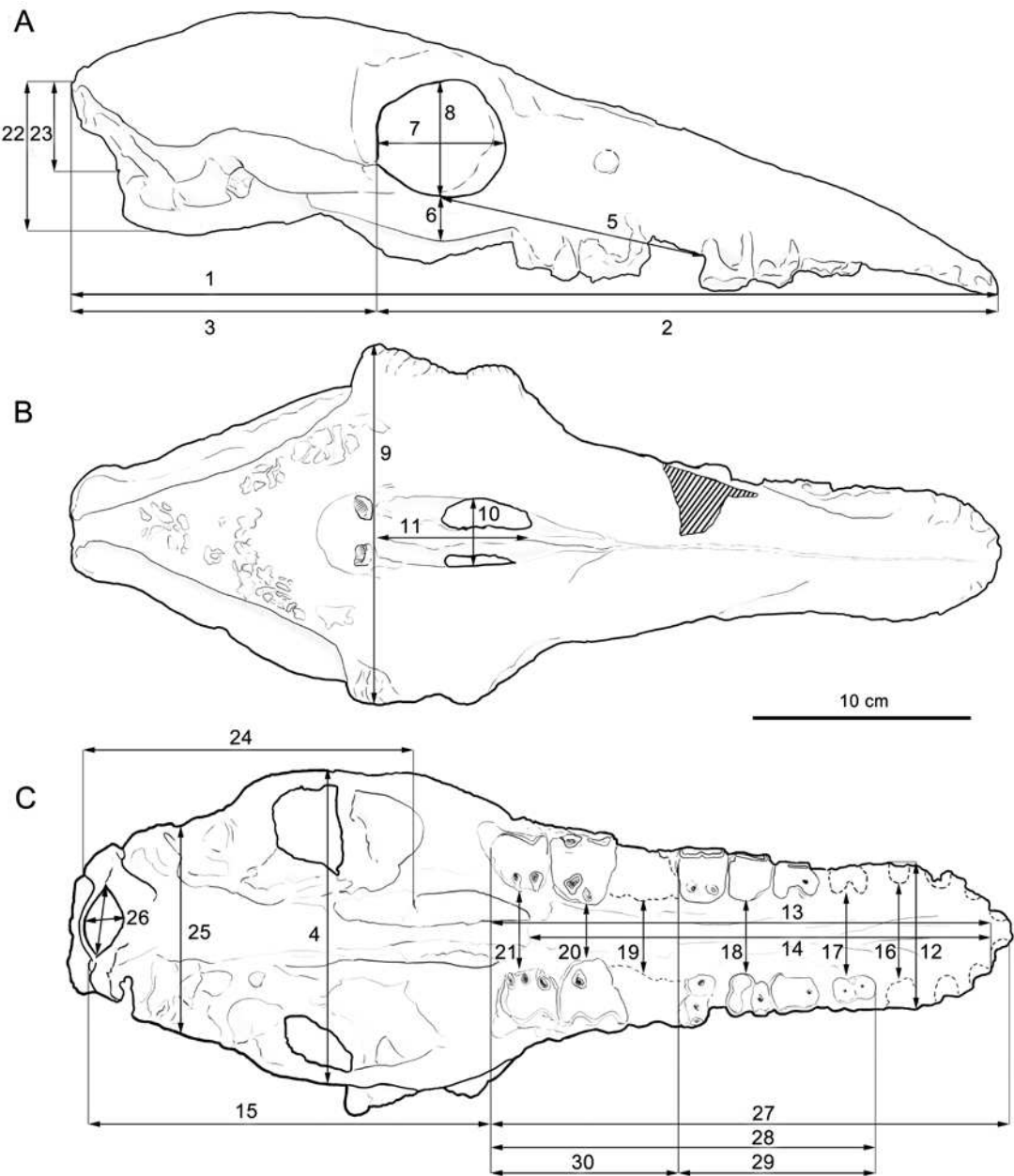


FIG. 4. Skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29 in **A**, right lateral, **B**, dorsal, and **C**, ventral views, with measurements (see table 2).

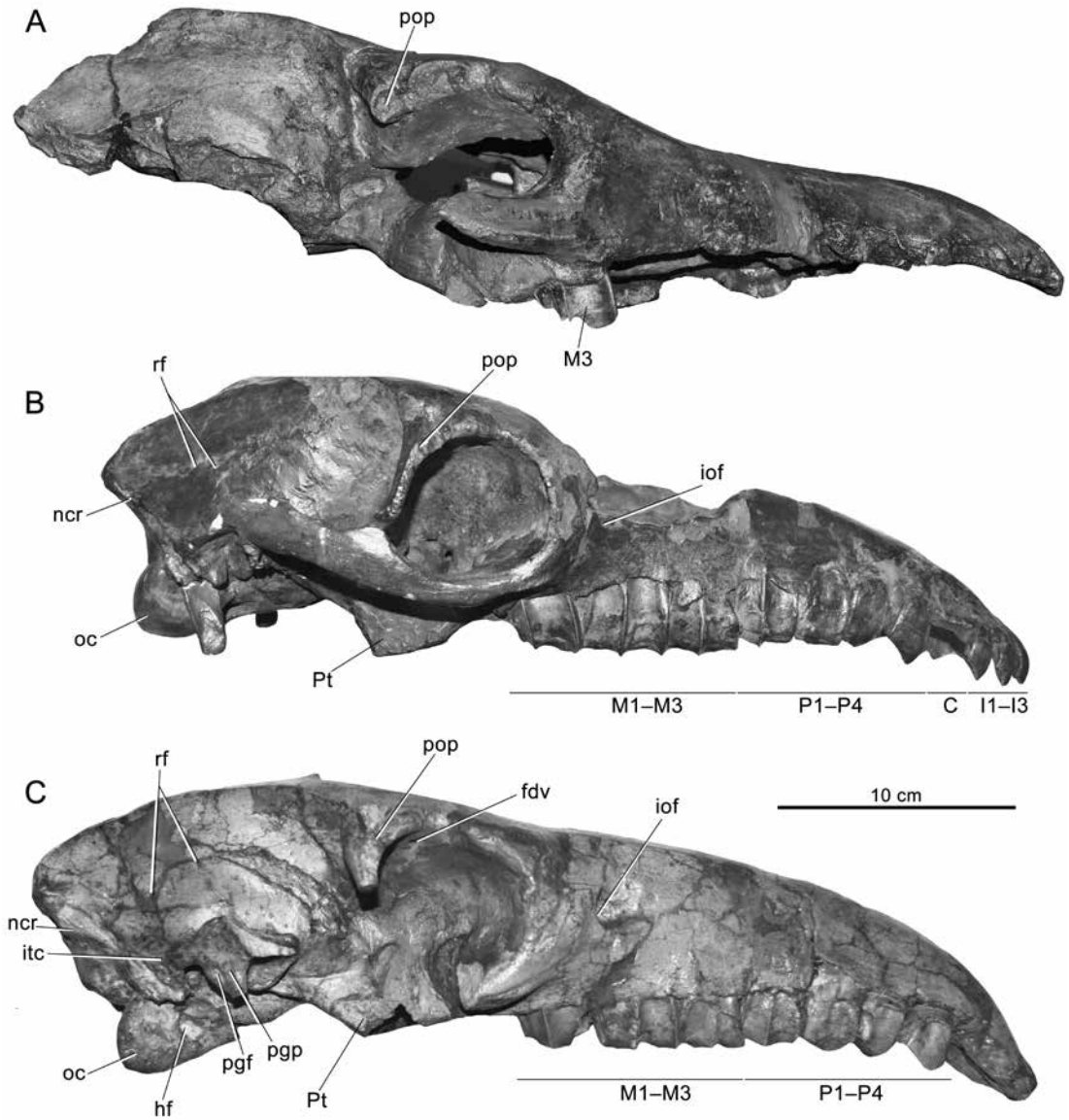


FIG. 5. Lateral views of skulls of **A**, *Huayqueriana cristata* MLP 37-III-7-2, Huayquerías Formation, Mendoza; **B**, *Promacrauchenia antiqua* MACN-PV 7986, Monte Hermoso Formation, Buenos Aires; and **C**, *Promacrauchenia calchaquiorum* MACN-PV 5528 (holotype), “estratos Araucanos,” Catamarca. Abbreviations: **C**, canine (alveolus); **fdv**, foramen for frontal diploic vein; **hf**, hypoglossal foramen; **I**, incisor; **iof**, infraorbital foramen; **itc**, infratemporal crest; **M**, molar; **ncr**, nuchal crest; **oc**, occipital condyle; **P**, premolar; **pgf**, postglenoid foramen; **pgp**, postglenoid process; **pop**, postorbital process of frontal; **Pt**, pterygoid; **rf**, foramina for rami temporales.

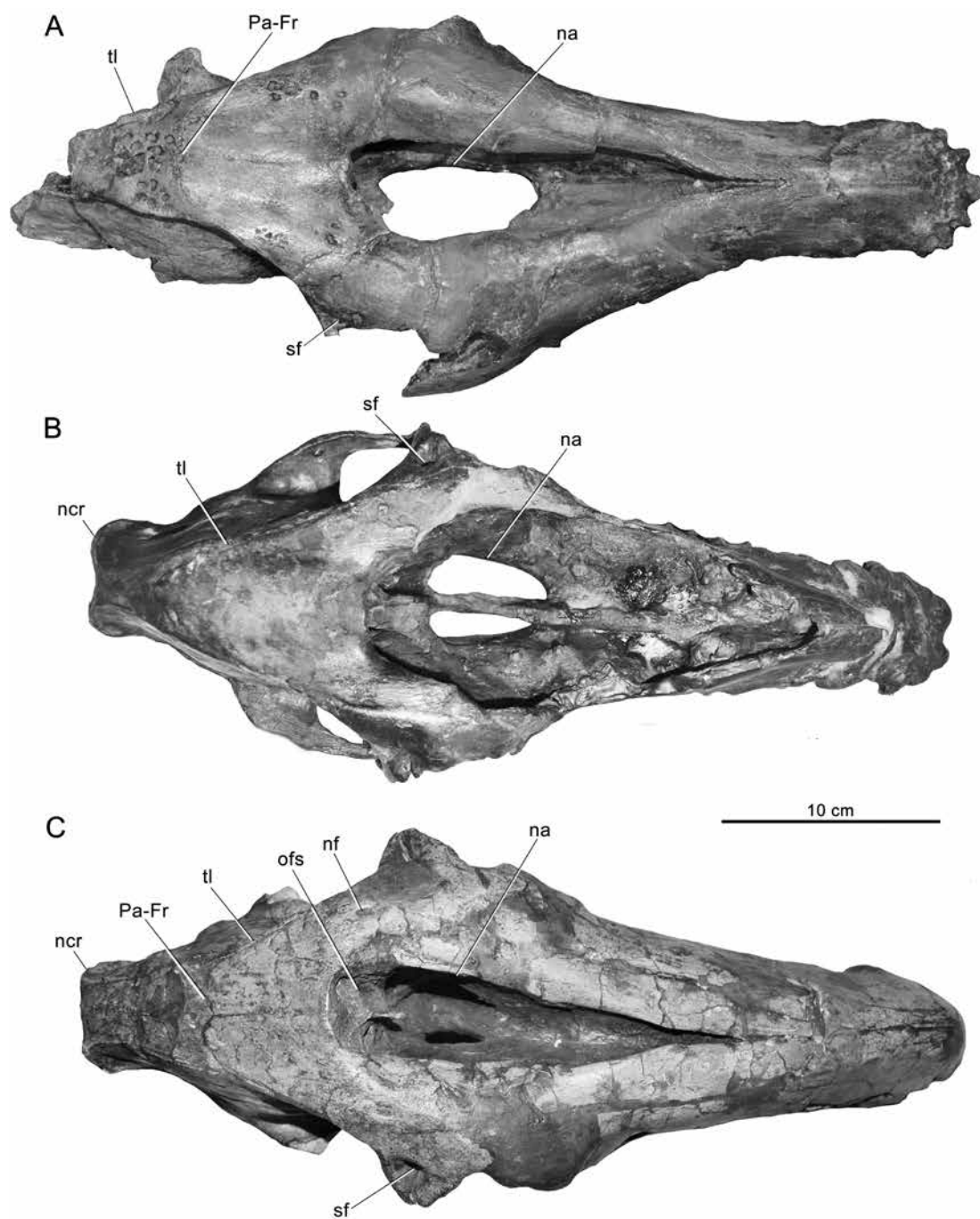


FIG. 6. Dorsal views of skulls of **A**, *Huayqueriana cristata* MLP 37-III-7-2; **B**, *Promacrauchenia antiquua* MACN-PV 7986; and **C**, *Promacrauchenia calchaquiorum* MACN-PV 5528 (holotype). For stratigraphic provenances, see figure 5. Abbreviations: **Fr**, frontal; **na**, nasal aperture; **ncr**, nuchal crest; **nf**, nutrient foramina; **ofs**, openings of frontal sinus; **Pa**, parietal; **sf**, supraorbital foramen; **tl**, temporal line.

3B), similar to conditions in *H. cristata* (Soria, 1986; fig. 6A) and *S. bravardi* MACN-PV 8903. In contrast, in *Pr. calchaquiorum* (fig. 6C) the maxillae join at the transverse level of M1, while in *M. patachonica* they meet at M2 (Fernández de Álvarez, 1940).

We infer that during ontogeny the rostral border of the nasal aperture migrated progressively caudally, as in the juvenile of *O. zeballosi* MHIN-UNSL-GEO-V 465a (fig. 8): the maxillae meet at the DP1/DP2 level, whereas in the adult of the same species MACN-PV 13671 they do not come into contact up to the level of, or even posterior to, the mesial portion of P3. Its precise position in MACN-PV 13671 cannot be fixed because the maxillary nasal border is incomplete caudal to the mesial part of P3.

Compared to conditions in most other mammals, the nasal aperture of macraucheniiines is remarkably derived, differing radically in its proportions and organization even from non-macraucheniiine macraucheniiids (Scott, 1910). Its derivative nature is especially obvious in the orientation of the passageway between the external opening of the nares and the nasopharynx. In most mammals this passageway tends to be coplanar with the palate, i.e., essentially horizontal in life position. In species with notably derived cranial modifications for head posture, such as humans, the passageway exhibits a sharp ventral bend at the level of the soft palate, such that the oropharynx joins the caudal nasal apertures at a near right angle (Hiatt and Gartner, 2001). In *Huayqueriana* and other macraucheniiines the entire passageway is normal to the palate (figs. 3A, C; 10, 11), with the result that the large oval choanae are directed dorsoventrally, not rostrocaudally. More remarkably still, and as revealed by CT scans, the rostralmost portion of the nasal cavity still retains its primitive position, occupying the snout almost to the tip (figs. 10, 11, 12C). However, it is now reduced to a diverticulum, lacking any kind of entrance above the incisor row and thus having no apparent primary role as a conduit for airflow. In the tapir, a similar space occupies the rostral part of the snout

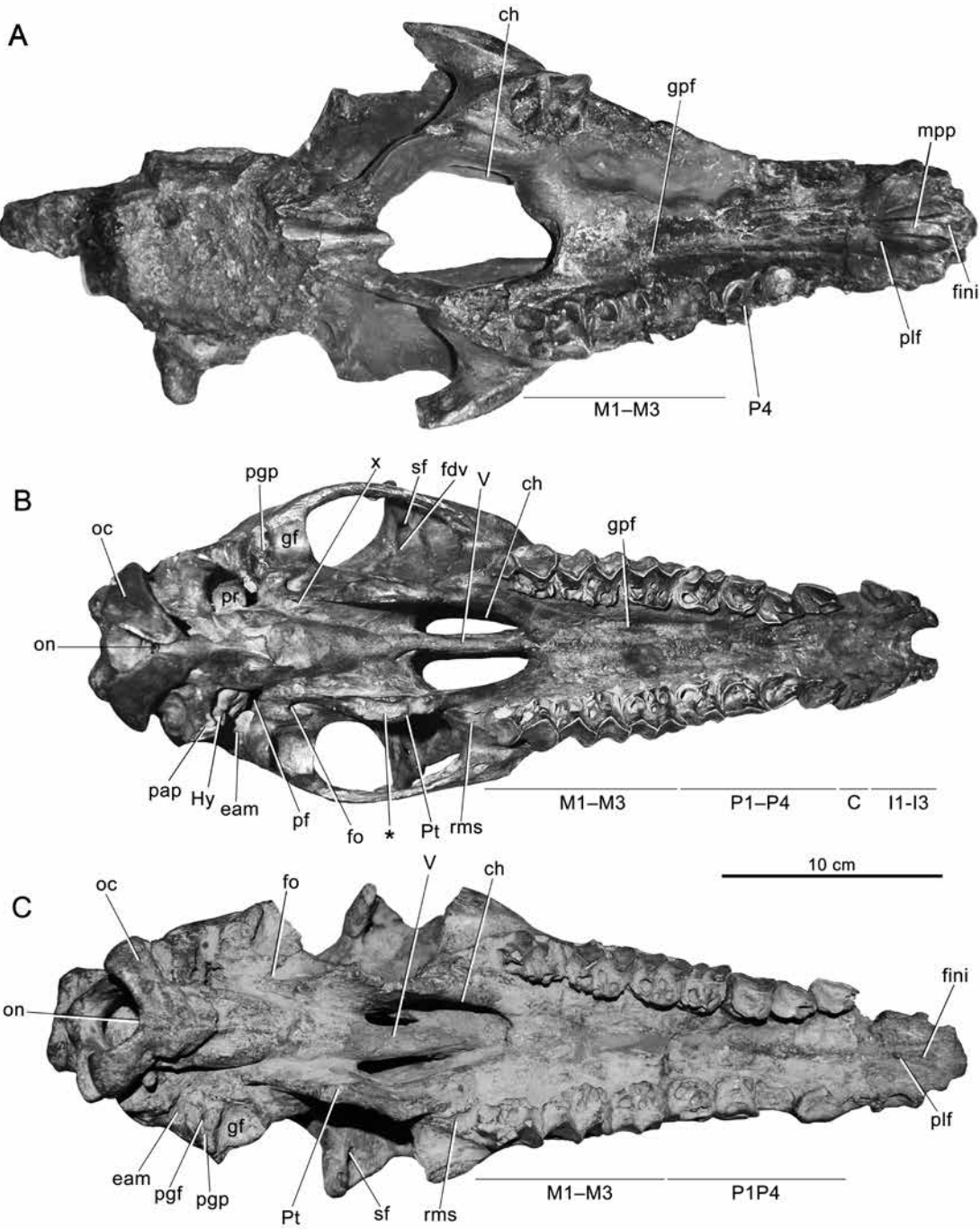
(hereafter referred to as meatus nasi ventralis, cf. Witmer et al., 1999).

In ventral view, the rostral border of the right choana in IANIGLA-PV 29 is located at the transverse level of the mesial border of M3, but the left is positioned slightly more caudally, its mesial border intersecting the middle of M3. This asymmetry is probably due to taphonomic distortion. In *S. bravardi* MACN-PV 8903, the rostral borders of both choanae are situated even with the mesial border of M3. In *Pr. antiquua* and *H. cristata* MLP 41-IV-29-4 they are positioned above the mesial half of M3; in *Pr. calchaquiorum*, above the midsection of the tooth; and in *H. cristata* (holotype), level with the distal border of the latter. In *O. zeballosi* (MACN-PV 13671), the choanae lie relatively further forward, along a plane running through the middle of M2.

In ventral view, the nasal septum presents as a thick, vertical lamina separating the large choanal apertures. Most of the bony septum is evidently formed by the hypertrophied vomer (figs. 11, 12F); on comparative grounds the perpendicular plate of the ethmoid must also be involved, although there is no evidence of a suture. Fernández de Álvarez (1940) established that in *M. patachonica* the dorsal margin of the vomer is sandwiched between the maxillae, such that its blade visibly protrudes into the external surface of the intermaxillary suture for virtually the entire length of the snout. In IANIGLA-PV 29, similar conditions are found, although the vomer's participation in the intermaxillary suture is more limited and the rostral end of the nasal septum appears to be formed exclusively by maxillary material (fig. 12D–F).

Unfortunately, delicate turbinal structures were partially destroyed during preparation of the choanae. However, enough remains of the ethmoturbinates were situated where expected, i.e., in relation to the caudodorsal end of the nasal cavity (figs. 10, 11, 13A), and not within the rostrally situated meatus nasi ventralis.

Although in IANIGLA-PV 29 there is no trace of sutures in the rear wall of the nasal aper-



ture, the bone immediately caudal to the nasal aperture is probably composed of the two nasal bones, indivisibly fused. In the past there has been much doubt regarding this character, in part because the region in question varies considerably in appearance in different macraucheniiines. Rusconi (1932) was under the impression that a similar blunt protuberance in *S. bravardi* (MACN-PV 13082; fig. 9B) was actually composed of frontal material, to which the nasals were loosely attached. Fernández de Álvarez (1940: 19) expressed a similar view in regard to *M. patachonica* (MACN-PV 2): “en la cresta anterior a la cavidad triangular del frontal, deben haber estado implantados los nasales. Estos faltan en el ejemplar, pero no deben haber sido muy desarrollados.” However, other authors (Burmeister, 1864; Soria, 1986), as well as Fernández de Álvarez (1940) in regard to non-*Macrauchenia* macraucheniiines, proposed that the nasals either participate in, or actually compose, the protuberance bordering on the caudal rim of the nasal aperture, in continuity with definite frontal material. More recently, Soria pointed out that in *H. cristata* (MLP 41-IV-29-4), the nasals “están reducidos y forman una ligera elevación redondeada, a modo de cresta, en el borde postero-medial de la abertura nasal” (Soria, 1986: 160).

Our reexamination of the relevant region in specimens studied by these authors, including MACN-PV 8903, MACN-PV 5528, MACN-PV 7986, MACN-PV 2, MLP 41-IV-29-4 (appendix 1), also failed to reveal any evidence of sutures. However, a juvenile assigned to *O. zeballosi* MHIN-UNSL-GEO-V 465a (Cerdeño et al.,

2008), in which several sutures and synchondroses are still partially patent, was found to exhibit the internasal suture and possibly part of the nasofrontal suture (fig. 8B). The nasofrontal suture may obliterate very early in ontogeny, as in elephants, in which it fuses sooner than the basioccipital-basisphenoid synchondrosis (e.g., *Loxodonta africana*) or immediately thereafter (e.g., *Elephas maximus*) (Rager et al., 2014).

The notion that nasals might actually be absent or detached in some specimens or species of macraucheniiines may have arisen from the peculiar condition of the dorsal surface of the bony nasal septum at the point at which it meets the caudal rim of the nasal aperture. To varying degrees in different macraucheniiine taxa, the last two or three centimeters of the septum broaden out into narrow wings defining a central depression. Often the surfaces of the wings are roughened, as though they supported a nonbony structure. To a degree the effect is similar to a suture, but in comparing various macraucheniiines to mammals having the usual form of the nasal skeleton, we have come to the conclusion that these wings must have supported the dorsocaudal terminus of the nasal septal cartilage. On positional grounds this makes sense, but it is harder to reconstruct the form that the cartilaginous nasal septum would have taken. Witmer et al. (1999) have shown that the septal cartilage in the tapir is a decidedly complicated structure that terminates rostrally in a long fibrous cord that arcs through or under the hydrostat (proboscis) to terminate in the tissues of the upper lip. Whether mac-

FIG. 7. Ventral views of skulls of **A**, *Huayqueriana cristata* MLP 37-III-7-2; **B**, *Promacrauchenia antiquua* MACN-PV 7986; and **C**, *Promacrauchenia calchaquiorum* MACN-PV 5528 (holotype). For stratigraphic provenances, see figure 5 caption. Abbreviations: **C**, canine; **ch**, choana; **eam**, external acoustic meatus; **fdv**, foramen for frontal diploic vein; **fini**, foramen interincisivum; **fo**, foramen ovale; **gf**, glenoid fossa; **gpf**, greater (= major) palatine foramen; **Hy**, hyoid (tympanohyal + stylohyal); **I**, incisor; **M**, molar; **mpp**, medial palatine process of premaxilla; **oc**, occipital condyle; **on**, odontoid notch; **P**, premolar; **pap**, paracondylar process of exoccipital; **pf**, piriform fenestra; **pgf**, postglenoid foramen; **pgp**, postglenoid process; **plf**, palatine fissure; **pr**, promontorium; **Pt**, pterygoid; **rms**, retromolar space; **sf**, supraorbital foramen; **V**, vomer. Letter **x**, identifies canal X for nerve of pterygoid canal and vascular structures (see text). Asterisk (*) identifies marked groove on ventral margin of pterygoid plate, especially evident on MACN-PV 7986.

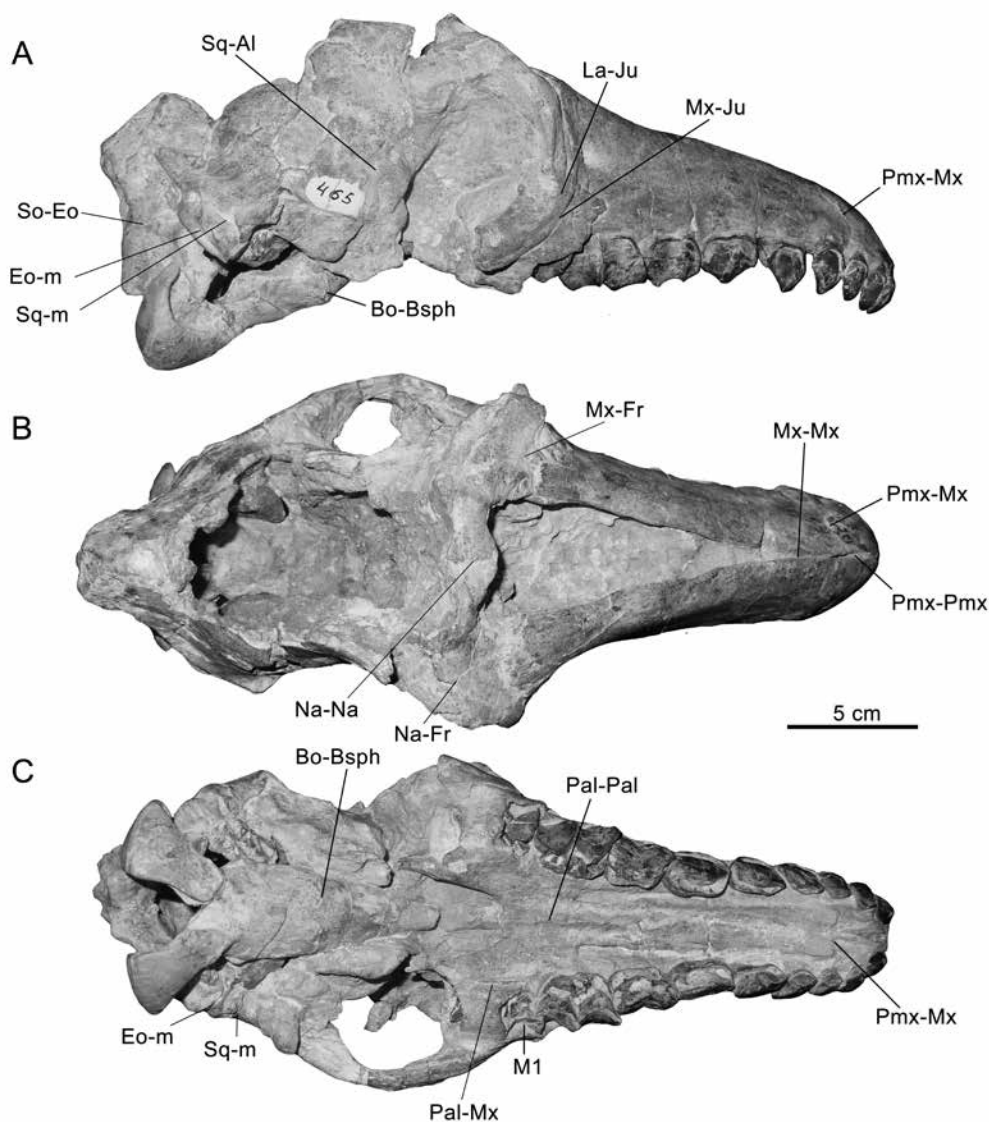


FIG. 8. Skull of *Oxydontherium zeballosi* MHIN-UNSL-GEO-V 465a from Río Quinto Formation, San Luis, in **A**, lateral, **B**, dorsal, and **C**, ventral views, with sutures indicated. Abbreviations: **Al**, alisphenoid; **Bo**, basioccipital; **Bsph**, basisphenoid; **Eo**, exoccipital; **Fr**, frontal; **Ju**, jugal; **La**, lacrimal; **Mx**, maxilla; **Na**, nasal; **Pal**, palatine; **m**, mastoid process of petrosal; **Pmx**, premaxilla; **So**, supraoccipital; **Sq**, squamosal.

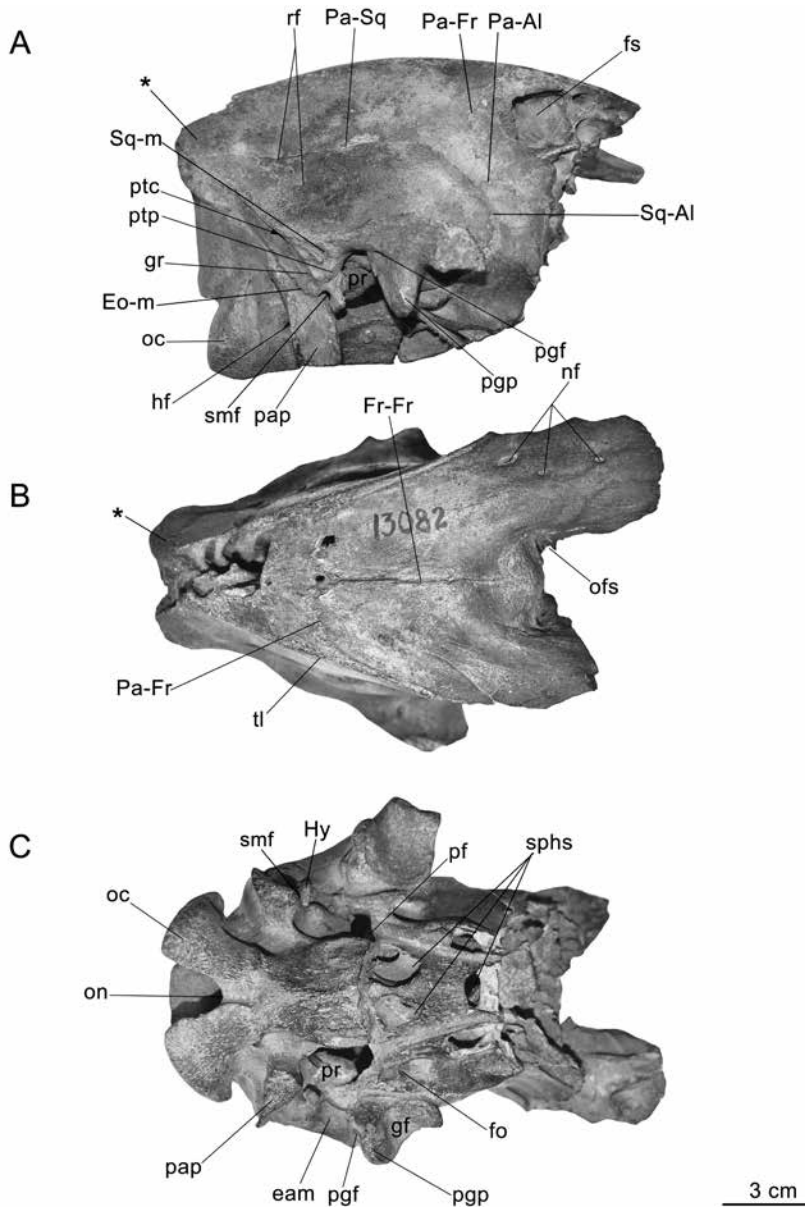


FIG. 9. Braincase of *Scalabrinitherium bravardi* MACN-PV 13082 from Ituzaingó Formation, Entre Ríos, in **A**, lateral, **B**, dorsal, and **C**, ventral views. Rostral part of specimen extensively damaged; apparent apertures on ventral aspect are not foramina but instead exposed parts of sphenoidal sinus (**sphs**). Other abbreviations: **Al**, alisphenoid; **eam**, external acoustic meatus; **Eo**, exoccipital; **fo**, foramen ovale; **Fr**, frontal; **fs**, frontal sinus; **gf**, glenoid fossa; **gr**, occipital groove; **hf**, hypoglossal foramen; **Hy**, hyoid (tympanohyal + stylohyal); **m**, mastoid process of petrosal; **nf**, nutrient foramina; **oc**, occipital condyle; **ofs**, openings of frontal sinus; **on**, odontoid notch; **Pa**, parietal; **pap**, paracondylar process of exoccipital; **pf**, piriform fenestra; **pgf**, postglenoid foramen; **pgp**, postglenoid process; **pr**, promontorium; **ptc**, posttemporal canal; **ptp**, posttympanic process of squamosal; **rf**, foramina for rami temporales; **smf**, stylomastoid foramen; **Sq**, squamosal; **tl**, temporal line. Asterisk (*) identifies suture-delimited element, probably interparietal.



FIG. 10. Virtual parasagittal section through choana of skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Light gray areas within skull represent unremoved matrix. Caudally-situated vacancy between nasal aperture and nasopharynx is here denoted as “true” nasal cavity (**tncav**), as it would have acted as the most direct air passageway in life. Primitive nasal cavity (**mnv**, meatus nasi ventralis) persists as a diverticulum, lacking an external aperture at front of rostrum. Note substantial thickness of rostralmost portions of premaxilla/maxilla at front of long, comparatively gracile rostrum. Other abbreviations: **chl**, cochlea; **crh**, position of cerebral hemispheres; cerebral cavity; **Ec**, ectotympanic; **fs**, frontal sinus; **mnv**, meatus nasi ventralis; **Mx**, maxilla; **ns?**, possible distorted portion of nasal septum; **pafs**, parietal extension of frontal sinus; **plf**, palatine fissure; **Pmx**, premaxilla; **sphs**, sphenoidal sinus; **t**, turbinate fragments in caudal portion of nasal cavity; **tncav**, true nasal cavity.

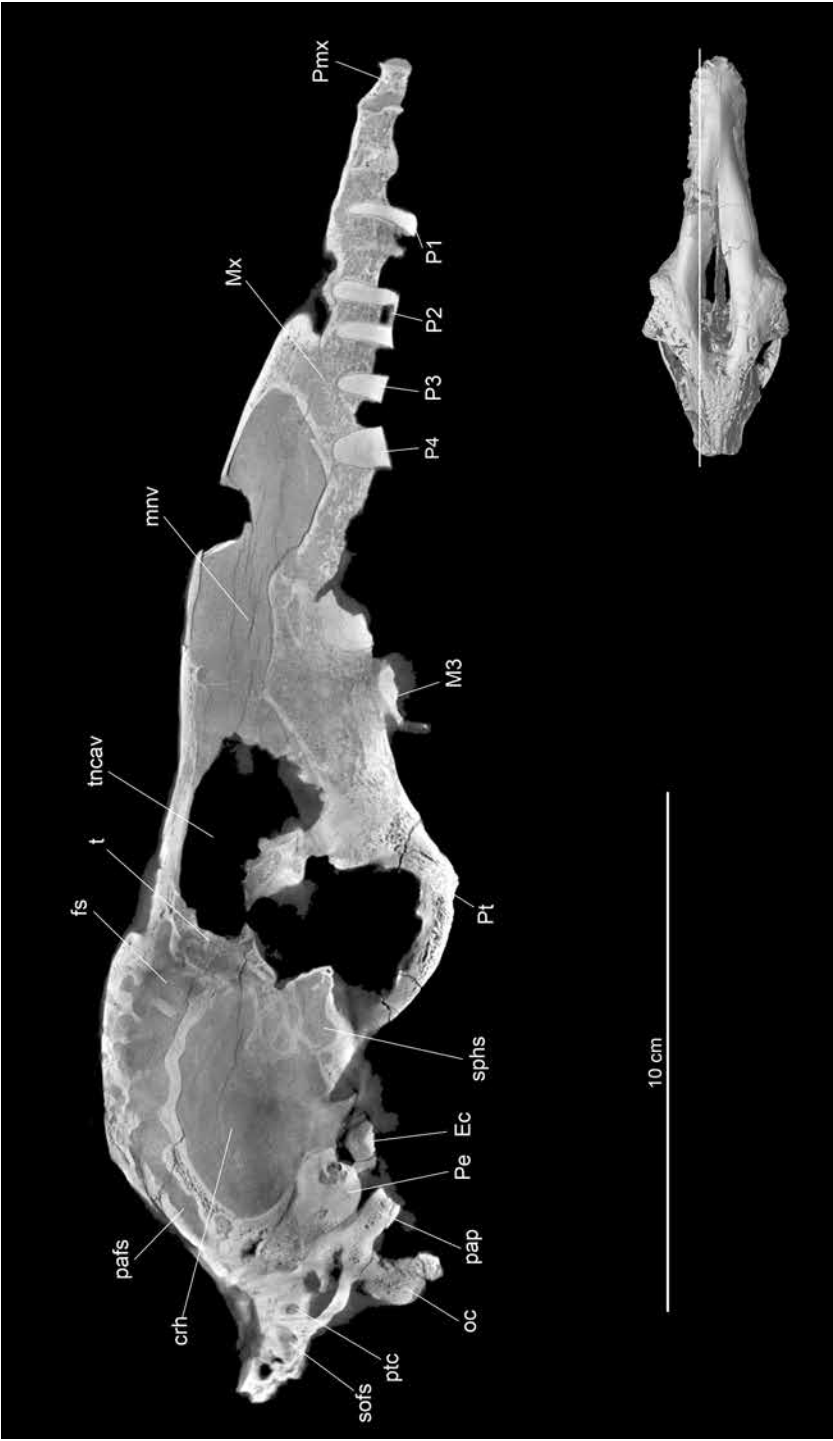


FIG. 11. Virtual parasagittal section through skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **crh**, position of cerebral hemispheres (cerebral cavity); **Ec**, ectotympanic; **fs**, frontal sinus; **Mx**, maxilla; **oc**, occipital condyle; **P**, premolar; **pafs**, parietal extension of frontal sinus; **pap**, paracondylar process of exoccipital; **Pe**, petrosal; **Pt**, pterygoid; **ptc**, posttemporal canal; **sofs**, supraoccipital extension of frontal sinus; **sphs**, sphenoidal sinus; **t**, turbinate fragments in caudal portion of nasal cavity; **tn cav**, true nasal cavity.

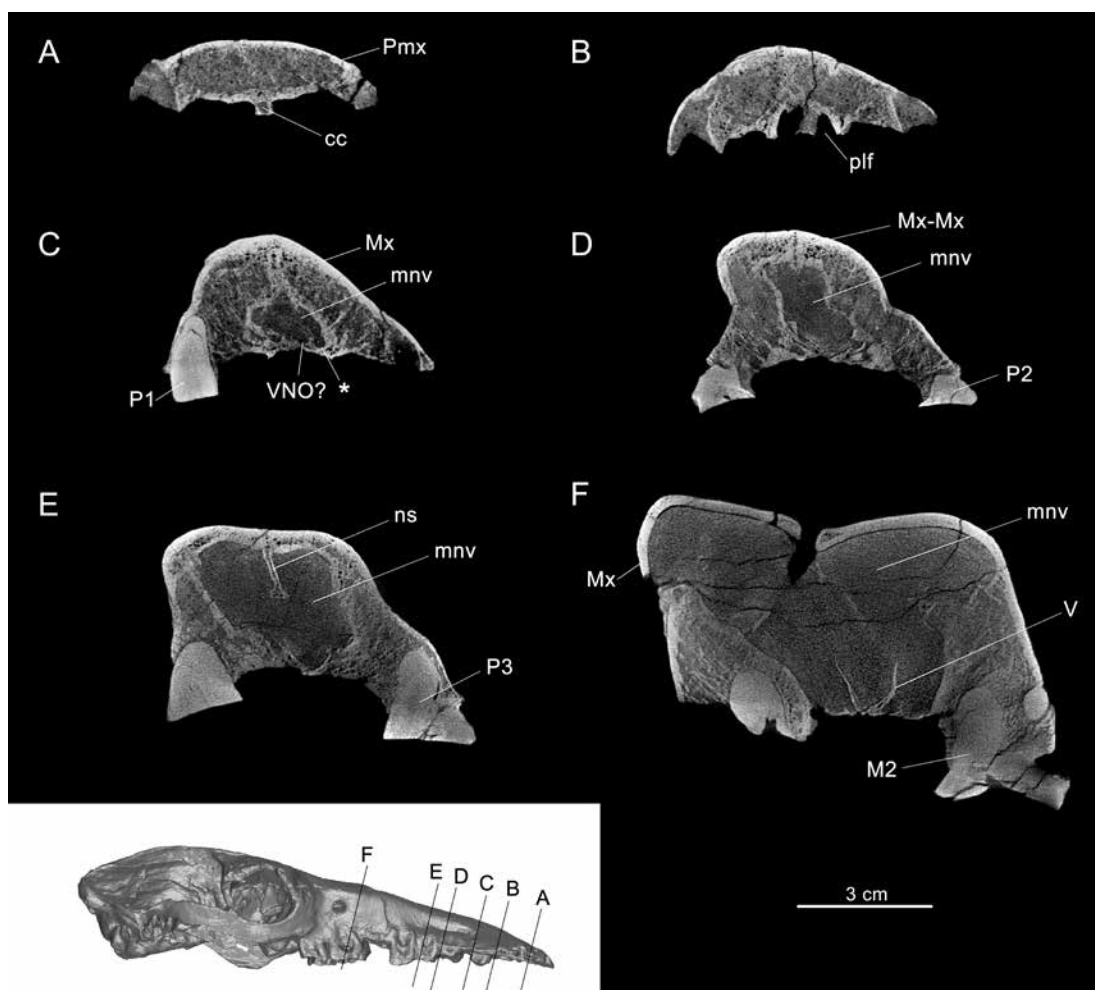


FIG. 12. A–F, Virtual coronal sections of skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **cc**, central canal; **M**, molar; **mnv**, meatus nasi ventralis; **Mx**, maxilla; **ns**, nasal septum; **P**, premolar; **plf**, palatine fissure; **Pmx**, premaxilla; **V**, vomer; **VNO?**, possible sulcus for vomeronasal organ. Asterisk (*) identifies sulcus for vascular structures (see text).

raucheniines possessed anything similar is of course unknown, although the unusual degree of vomer exposure in the intermaxillary suture may imply actual prolongation of the nasal cartilage onto (or into) the face.

In summary, the preponderance of evidence indicates that the nasals were very small in macraucheniines and projected little or not at all beyond the caudal rim of the nasal aperture, a decided similarity to elephants, less so to other taxa possessing prominent nasal ornamentations such as *Tapirus* and *Saiga*.

PALATE

In ventral view (fig. 3C), the hard palate of IANIGLA-PV 29 is narrow and uniform in width (table 2; fig. 4C). Allowing for size, the palate appears narrower than in other macraucheniines (fig. 7), although this could be a consequence of taphonomic compression.

The rostralmost part of the palate exhibits a midline aperture and an associated canal, as well as two laterally positioned, sharply bordered dehiscences, the palatine fissures (= palatal

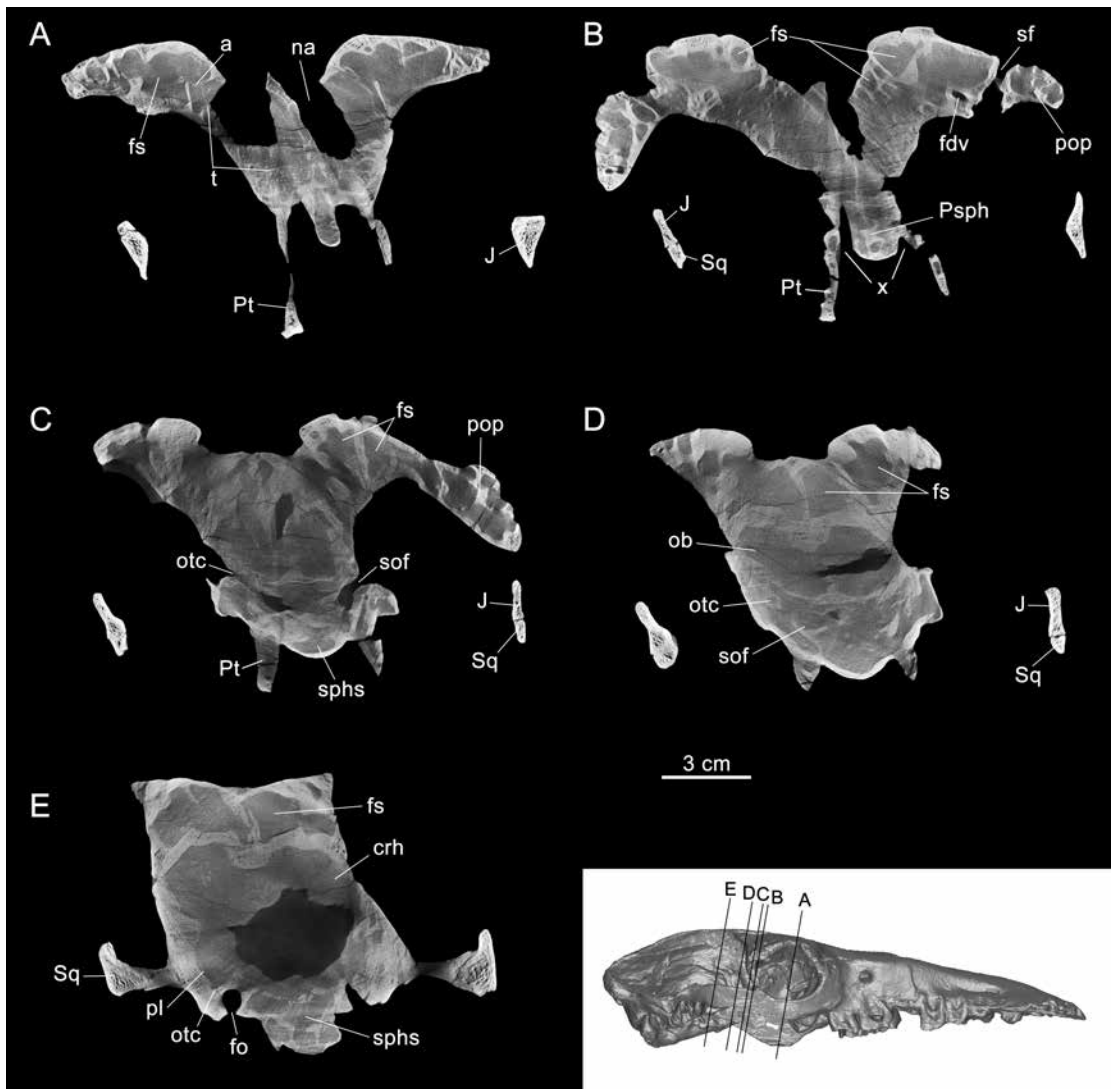


FIG. 13. A–E, Virtual coronal sections of skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **a**, aditus to frontal sinus; **crh**, position of cerebral hemispheres; **fdv**, foramen for frontal diploëic vein; **fo**, foramen ovale; **fs**, frontal sinus; **J**, jugal; **na**, nasal aperture; **ob**, position of olfactory bulb; **otc**, orbitotemporal canal; **pl**, position of piriform lobe; **pop**, postorbital process of frontal; **Psph**, presphenoid; **Pt**, pterygoid; **sf**, supraorbital foramen with perforating canal; **sof**, sphenoorbital fissure; **Sq**, squamosal (zygomatic process); **sphs**, sphenoidal sinus; **t**, turbinals. Letter **x**, identifies canal X for nerve of pterygoid canal and vascular structures (see text).

fenestrae) (fig. 3C). The midline aperture, which we consider the operational homolog of the perissodactyl foramen interincisivum (= foramen palatinum medium; Maier, 2002), is located at the rostral end of a long (3 cm), somewhat elevated tube (hereafter, the central canal; fig. 12A). This foramen is also present in *Huayqueri-*

ana cristata MLP 41-IV-29-4. Unlike the primitive incisive foramina of mammalian anatomy, foramen interincisivum is unpaired.

The raised ridge housing the central canal probably exists in other macrauchenini (e.g., *O. zeballosi* MHIN-UNSL-GEO-V 465a; *M. patachonica* MACN-PV 2), but its identity as a conduit is obvi-

ous only when scans are examined. The foramen interincisivum is broad and V-shaped in ventral view, and is situated just posterior to the alveoli of the mesial incisors, as in other macraucheniniines (e.g., *S. bravardi* MACN-A 1270, *Pr. calchaquiorum*, and *O. zeballosi* MHIN-UNSL-GEO-V 465a; figs. 7, 8C). This opening rapidly narrows into the central canal proper, which is only a few millimeters wide (fig. 12A). The canal becomes continuous with the meatus nasi ventralis at the approximate level of the rostral border of P1.

Two deep palatal grooves are developed lateral and caudal to the central canal (fig. 3C), as in other macraucheniniines (figs. 7, 8). These grooves open into the palatine fissures at the level of the canine alveoli (figs. 10, 12B; contra Scott 1910: 117, who thought only grooves were present). These fissures are caudally continuous with grooves on the floor of the meatus nasi ventralis, possibly for divisions of the greater (= major) palatine artery and nerve. Thus, the fissures presumably reflect the passage of those soft tissue structures from the meatus nasi ventralis, into the rostral part of the upper jaw.

There are at least two possibilities for the contents of the central canal, based on conditions in modern perissodactyls (*Equus*). The first concerns the arteria incisiva (= palatolabial artery), a terminal division of the greater palatine artery (Sisson and Grossman, 1975; Vogt, 2011). The arteria incisiva is formed by the union of two branches of the greater palatine artery to form a single median vessel, which enters the foramen interincisivum immediately behind the medial incisors (Sisson and Grossman, 1975). The vessel thereafter ascends dorsocaudally to emerge from its exit foramen situated on the nasal sill. In *Equus*, the arteria incisiva is transmitted via the foramen interincisivum to the upper lip, where it releases dorsal and ventral septal branches destined for the nasal cavity and the transversus nasi muscle and upper lip. By contrast, *Huayqueriana*, like *Tapirus*, lacks a large foramen opening dorsally on the forward margin of the rostrum. Thus, if the central canal contained the homolog of the arteria incisiva of the horse, it differed from the latter in its area of supply.

The second possibility for the content of the central canal is related to innervation. Although in mammals the upper lip is mainly supplied with sensory fibers by the infraorbital nerve, applied anaesthesia on the horse (Vogt, 2011) indicates that the nasopalatine nerve also passes through the foramen interincisivum. In most mammals, the nasopalatine nerve, a long branch of the posterior superior nasal branches of CN V₂, runs between the nasal septum and the mucus membrane to reach ports within the incisive canal, through which it passes to communicate with the greater palatine nerve (Hiatt and Gartner, 2001). In shrews (e.g., *Neomys*) the nasopalatine nerve crosses the hard palate via a single midline opening in the rostral part of the snout (Maier, 2002), which strongly recalls the condition in *Huayqueriana*.

We consider it highly unlikely that the nasopalatine (= incisive) ducts, the structures that connect the vomeronasal organ (VNO) (= Jacobson's organ) to the external environment, passed through the foramen interincisivum/central canal complex rather than the palatal fissures. Although probably present in life, the VNO's position in the fossil is uncertain. As may be seen in figure 12C, small paired depressions in the rostralmost part of the meatus nasi ventralis are consistent with VNO presence, but whether they are actually related to this organ or simply mark the passage of blood vessels, alveolar nerves, or attachment points of the septal cartilage cannot be settled. In any case, as seen in the horse and tapir (von Mering, 1994; Witmer et al., 1999), for example, the nasopalatine ducts are paired and follow separate, oblique pathways to the ipsilateral VNO. The midsagittal position of the central canal is inconsistent with such a routing. It is much more likely that the ducts passed through the wide palatine fissures, as in modern perissodactyls (e.g., von Mering, 1994; Witmer et al., 1999).

Caudal to the foregoing, the hard palate is incised by two major grooves that run in parallel from the greater palatine foramina. The greater palatine foramina are small and asymmetrically

located: the one on the right is at the level of the M2/M3 embrasure, while that on the left is in line with the middle of M3 (fig. 3C). There is variation in the position of the palatine foramina in our comparative set. In *O. zeballosi* MACN-PV 13671, the right aperture is situated at the level of the distal portion of M1; in *Pr. antiquua* (fig. 7B) the foramina intersect a line drawn through the midpart of M1 (similar to *H. cristata* MLP 41-IV-29-4 and *M. patachonica* MACN-PV 2; Fernández de Álvarez, 1940). In *S. bravardi* MACN-PV 8903 the palatine foramina open at the level of the mesial half of M2, while in *Pr. calchaquiorum* (fig. 7C) and *X. bahiense* they open in line with the middle of M2 (Cartelle and Lessa, 1988: 6). In the holotype of *H. cristata* (fig. 2B), the right greater palatine foramen opens at the level of the middle of M2 while the left does so at the level of the mesial portion of M3.

The retromolar space on the maxilla of IANIGLA-PV 29 is almost as long as the M3 itself (fig. 3C), as in other examined macraucheniiines (fig. 7), except for *O. zeballosi* MACN-PV 13671 and MACN-PV 17745 in which this space is somewhat shorter.

The pterygoids of IANIGLA-PV 29 are roughly triangular and platelike in shape. Even though we are aware that the pterygoid hamulus is a delicate structure often broken in skulls, our observations on several macraucheniiine specimens strongly indicate that there is no hamulus in *Huayqueriana* (fig. 3C), a fact that Scott (1910) previously noted for *Theosodon*. However, in *Huayqueriana* the ventral margin of the pterygoid plate is of unusual construction. Here, a curved longitudinal groove indents the margin along its entire length. A similar feature is seen in *Pr. antiquua* (fig. 7B) and *Promacrauchenia* sp. MLP 29-X-10-16, in which the groove is deep enough to partially divide the pterygoid plate into inner and outer moities. Its smooth floor and distinctive form are consistent with its occupation in life by a muscle belly or tendon (?tensor veli palatini), suggesting that this feature may have acted as a pulley to tense soft tissue structures related to the choanal opening.

ORBIT

Because of the extraordinary length of the rostrum, the orbits seem to be situated relatively caudally, as in other macraucheniiids. Positional relationships of the molar teeth to the orbit vary in our comparative set. In IANIGLA-PV 29 (fig. 3A) and in the holotype of *H. cristata* (fig. 2A), the rostral border of the orbit is almost level with the distal border of M3, similar to *Pr. calchaquiorum* (fig. 5C). In *Pr. antiquua* (fig. 5B), *S. bravardi* MACN-PV 8903, and *H. cristata* MLP 41-IV-29-4 (fig. 4A), the border is level with the middle of M3, while in *O. zeballosi* it intersects a plane drawn through the mesial portion of M3. In *M. patachonica*, the orbital margin lies behind the transverse position of the last molar (Fernández de Álvarez, 1940).

The orbital borders are prominent (fig. 3B) and feature a massive, ventrally projecting post-orbital process (fig. 3A). The process does not form a complete, *Equus*-like postorbital bar because there is a small gap between the zygomatic process of the frontal and that of the squamosal, similar to the condition found in *Pr. antiquua* and *Pr. calchaquiorum*. In *Theosodon* (Scott, 1910) and *Cramauchenia* (Dozo and Vera, 2010), the gap is relatively larger. By contrast, in *X. bahiense* and *M. patachonica* the bar is continuous and thus the orbit is completely encircled by bone (Scott, 1910; Soria, 1986; Cartelle and Lessa, 1988). According to Rovereto (1914, plate XXIX.4), the postorbital bar was reportedly complete in the holotype of *H. cristata*. This part of the skull is unfortunately now missing (fig. 2) and the specimen is in any case clearly dorsoventrally compressed. Considering how narrow the gap is in IANIGLA-PV 29 and *Pr. antiquua*, the apparent absence of a gap in the holotype of *H. cristata* may likewise be an artifact of taphonomic deformation. As the actual situation is ambiguous, we prefer to consider complete closure as uncertain in the holotype. There is a post-orbital constriction in IANIGLA-PV 29 as in other macraucheniiines, although this is not apparent in dorsal view because of the lateral projection of the temporal lines.

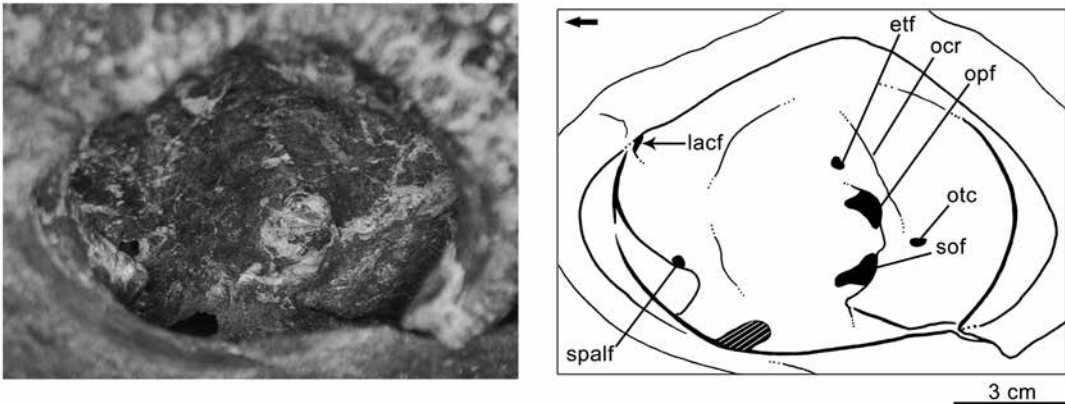


FIG. 14. Detail of orbit of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **etf**, ethmoidal foramen; **lacf**, lacrimal foramen; **ocr**, orbital crest; **opf**, optic foramen; **otc**, orbitotemporal canal; **sof**, sphenoorbital fissure; **spalff**, sphenopalatine foramen. Arrow points rostrally.

There is a small, single lacrimal foramen opening inside the orbit (fig. 14), as in *Pr. calchaquiorum* and *M. patachonica*. Each orbital roof is pierced by two foramina (figs. 3B, 13B). As seen from within the orbit, the lateral aperture opens on the summit of the postorbital process of the frontal (fig. 13B). We interpret this channel as the supraorbital foramen for the supraorbital artery and its accompanying vein. In adult mammals the artery is usually a branch of the deep temporal or maxillary artery, supplying the muscles and skin above the orbit, as in the horse (Sisson and Grossman, 1975). The medial aperture passes into the substance of the bone of the medial wall of the orbit (fig. 13B). We infer that this passageway carried a vein only, as seen in certain extant mammals (Wible and Gaudin, 2004; Wible, 2008), in this case draining the frontal diploë to the supraorbital vein. Similar foramina can be found on the orbital roof of other macraucheniniines (*Pr. antiquua*, *Pr. calchaquiorum*, and *M. patachonica*) (figs. 5–7).

The orbitotemporal fossa of IANIGLA-PV 29 (fig. 14) is very similar to that of some notoungulates (e.g., Toxodontia; Gabbert, 2004; Forasiepi et al., 2015b). There are three main apertures, dorsally bordered by the orbital crest. The two larger openings—the sphenoorbital fissure and the optic foramen—lie close to each other on the floor of the fossa. The remaining

aperture—the ethmoidal foramen—is much smaller and lies rostradorsally.

The sphenopalatine foramen (fig. 14), level with and rostral to the sphenoorbital fissure. This opening is large, oval, and communicates with the nasal cavity. On the left side of the skull, there is an opening at the base of the orbital crest, which suggests the presence of a small vessel communicating with the orbital cavity, here interpreted as the orbitotemporal canal (figs. 13C, 14; see Cerebral Endocast). A similar small opening in the orbital crest, at the transverse level of the optic foramen, was bilaterally present in *M. patachonica* (MACN-PV 2).

Vault

The skull roof is roughly triangular in dorsal view, ornamented by a network of irregular pits and scars of uncertain significance (fig. 3B). This roughened surface continues onto the postorbital process and supraorbital area. There is a small nutrient foramen on each side of the skull at the level of the caudal border of the nasal aperture. Similar openings were observed on the vault of *Pr. antiquua*, *Pr. calchaquiorum*, *M. patachonica* MACN-PV 2, and *S. bravardi* MACN-PV 13082 and MACN-PV 8903. The frontal is flat in *Pr. antiquua* (fig. 6B), *Pr. calchaquiorum* (fig. 6C),

and *S. bravardi* (fig. 9B); *M. patachonica* and *W. delacroixi* differ in that their frontals exhibit a number of large, shallow fossae grouped immediately caudal to the nasal aperture (Burmeister, 1864; Rusconi, 1932; Fernández de Álvarez, 1940). In *M. patachonica* MACN-PV 2 these fossae, partially subdivided by low septa, occupy a reniform space that is more than 7 cm wide and at least 3 cm deep. Interestingly, *Tapirus* displays differently positioned fossae for the so-called nasofrontal sacs originally described by Murie (1872) (= diverticulum meatus of Witmer et al., 1999). In the case of the tapir, the fossae contain glandular tissues that are connected by a channel to the nasal cavity (Witmer et al., 1999).

The broadly curving temporal lines on the rear part of the vault are very distinct and converge in the direction of the nuchal crest, which they join separately (fig. 3B). Despite their proximity caudally, the temporal lines do not actually meet in the midline and consequently there is no sagittal crest in IANIGLA-PV 29. There is instead a flat polygonal area, seen also in MLP 41-IV-29-4, a specimen referred to *H. cristata* (fig. 6A), *Pr. antiquua* (fig. 6B), *Pr. calchaquorum* (fig. 6C), *S. bravardi* (fig. 9B), *M. patachonica*, and, apparently, in *X. bahiense* (Burmeister, 1864; Rusconi, 1932; Fernández de Álvarez, 1940; Soria, 1986; Cartelle and Lessa, 1988). The holotype of *H. cristata* (fig. 2C), by contrast, possesses a short, caudally situated sagittal crest, resembling non-macraucheniine litopterns in this regard (Scott, 1910; Soria, 1986; Dozo and Vera, 2010). This difference may correspond to intraspecific variation (see Discussion). The nuchal crests of IANIGLA-PV 29 are also substantial, and connect with a sharply defined infratemporal crests about the level of the external acoustic meatus, directed toward the zygoma on each side of the skull.

The attachment area for the temporalis muscle is likewise bordered by three prominent crests: the temporal line dorsally, nuchal crest caudally, and the infratemporal crest ventrally. There are several foramina in the caudal part of the temporal fossa, some of which are associated with grooves marking the passage of vessels. Similar openings are present in several other litopterns and have been

given names such as foramen postsquamosale or foramen postparietale by different authors (e.g., in *Thoatherium minusculum*, *Tetramerorhinus lucarius*, *Cramauchenia normalis*, species of *Theosodon*, *H. cristata*, *Promacrauchenia* spp., *S. bravardi*, *M. patachonica*; Rusconi, 1932; Scott, 1910; Dozo and Vera, 2010; Fernández de Álvarez, 1940; figs. 2C, 6, 8A). We interpret these openings as conforming to foramina for rami temporales, seen in many other mammals (see Cerebral Endocast) and recently described for certain notoungulates (MacPhee, 2014; Forasiepi et al., 2015b; see also squamosal-parietal fenestration in García-López and Powell, 2011: fig. 1C).

ZYGOMATIC ARCH

The short and slender zygomatic arch is formed by the jugal and squamosal, which define between them an obliquely oriented suture (fig. 3A). The maxilla does not contribute to the ventral orbital rim in lateral view. The suture between the maxilla and jugal is almost vertical and above the M3 (this suture is oblique in the juvenile specimen of *O. zeballosi* MHIN-UNSL-GEO-V 465a; fig. 8A).

The zygomatic process of the squamosal is lengthy and extends onto the caudoventral margin of the orbit, as in other macraucheniids (e.g., *Cr. normalis*, *Pr. antiquua*, *M. patachonica*). The squamosal has a small, triangular frontal (= dorsal) process that contributes to the osseous caudal border of the orbit and thereby provides attachment for the postorbital ligament separating the orbital and temporal fossae (Wible, 2003). In other mammals this process is normally formed by the jugal (Wible, 2003; Sisson and Grossman, 1975), which, in conjunction with the postorbital process of the frontal, form the caudal border of the orbit.

BASICRANIUM

The basicranium in IANIGLA-PV 29 is long and narrow (fig. 15), as in other macraucheniids (fig. 7), with a convex ventral surface. Scott (1910) described a longitudinal crest on the basioccipital of *Theosodon* and observed that it

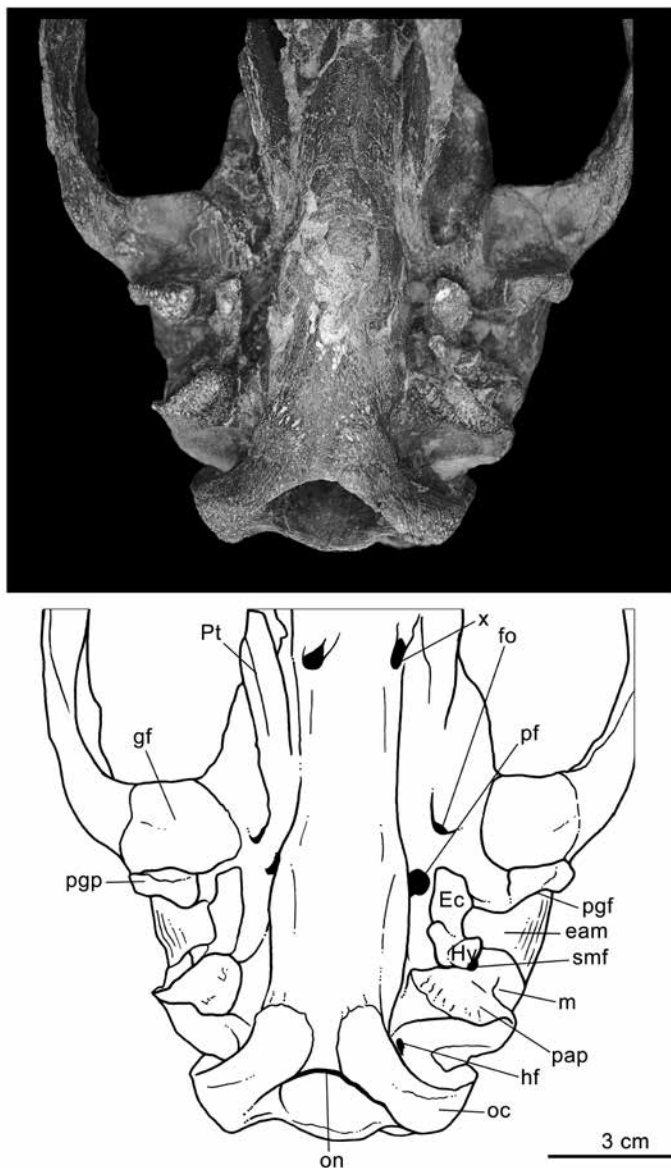


FIG. 15. Posterior portion of skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, in ventral view. Abbreviations: **eam**, external acoustic meatus; **Ec**, ectotympanic; **fo**, foramen ovale; **gf**, glenoid fossa; **hf**, hypoglossal foramen; **Hy**, hyoid (tympanohyal + stylohyal); **m**, mastoid process of petrosal; **oc**, occipital condyle; **on**, odontoid notch; **pap**, paracondylar process of exoccipital; **pf**, piriform fenestra; **pgf**, postglenoid foramen; **pgp**, postglenoid process; **Pt**, pterygoid; **smf**, stylomastoid foramen. Letter **x** identifies canal X for nerve of pterygoid canal and vascular structures (see text). Faintly visible on the specimen's left side is the large sulcus running between the piriform fenestra and canal X.

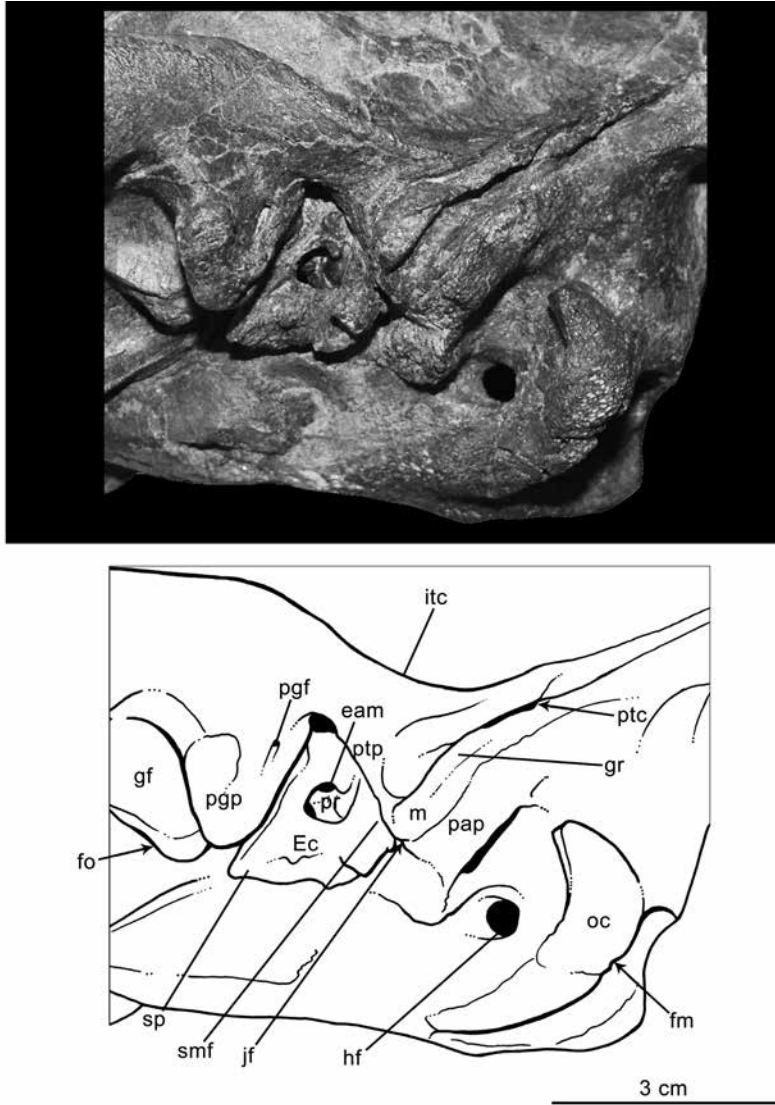


FIG. 16. Posterior portion of skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, in left ventrolateral view. Abbreviations: **eam**, external acoustic meatus; **Ec**, ectotympanic; **fm**, foramen magnum; **fo**, to foramen ovale; **gf**, glenoid fossa; **gr**, occipital groove; **hf**, hypoglossal foramen; **itc**, infratemporal crest; **jf**, to jugular foramen; **m**, mastoid process of petrosal; **oc**, occipital condyle; **pap**, paracondylar process of exoccipital; **pgf**, postglenoid foramen; **ppg**, postglenoid process; **pr**, promontorium; **ptc**, posttemporal canal; **ptp**, posttympanic process of squamosal; **smf**, stylomastoid foramen; **sp**, styliform process of ectotympanic.

became more pronounced with age. In the examined macraucheniines, a longitudinal crest is present in *Pr. antiquua*, *Pr. calchaquiorum*, and *S. bravardi* (figs. 7B, C, 9C). Despite the advanced ontogenetic stage of IANIGLA-PV 29, this feature is either negligible or absent.

The glenoid fossa is small and nearly circular. The postglenoid (= retroarticular) process is slightly wider than tall and pierced by a large postglenoid (= retroarticular) foramen as seen in posterior view (figs. 15, 16). The posttympanic (= retrotympanic) process of the squamosal is short and appressed to the mastoid process of the petrosal as in *Theosodon* (Scott, 1910), *M. patachonica*, *S. bravardi* (= processus retrozygomaticus of Rusconi, 1932), *O. zeballosi* (MHIN-UNSL-GEO-V 465a), and *Pr. calchaquiorum*. The posttympanic process is separated by a deep sulcus (see below) from the occipital paracondylar process (= paraoccipital process of Scott, 1910; Rusconi, 1932; Fernández de Álvarez, 1940).

The wide foramen ovale (figs. 13E, 15, 16) lies medial to the glenoid cavity and faces forward and downward. This opening has been previously identified in other macraucheniids as the foramen rotundum (Rusconi, 1932) or a confluent foramen ovale/rotundum (Scott, 1910). From examination of the endocast, we conclude that the only nerve piercing this opening was CN V₃, in agreement with observations by Simpson (1933) and Fernández de Álvarez (1940) for other macraucheniines. *Huayqueriana* and its allies studied to date evidently lacked a separate foramen rotundum (see also Cerebral Endocast) and the foramen rotundum was confluent with the speno-orbital foramen.

On either side of the interpterygoid space on the ventral surface of IANIGLA-PV 29, there are two prominent features. One is a pronounced groove running from the piriform fenestra along the medial side of the pterygoid process to a large canal (canal X; asterisk in fig. 15). Canal X perforates the lateral margin of the central stem close to the original location of the basisphenoid/pre-sphenoid synchondrosis. As large apertures situ-

ated in or near the mesocranial portion of the central stem are rare or absent in other mammals, we at first thought they were simply artificial dehiscences into the large pneumatic cells penetrating the sphenoid complex. Comparison with other macraucheniid specimens convinces us that they are original, although we have not been able to fully resolve what they carried, as the following discussion illustrates.

In IANIGLA-PV 29 (figs. 13B, 17B-D), canal X is more or less vertically oriented and apparently opened within the floor of the orbit. Because of preparation artifacts as well as the condition of the orbital areas in the specimen, the precise position of the dorsal ports for these canals is uncertain. Similar openings are seen in *Pr. antiquua* MACN-PV 7986 (fig. 7B), but they are situated much more caudally, at the level of foramen ovale, where they penetrate the pterygoid processes. From the morphological standpoint they are similar, however, in that they run within the interpterygoid or pharyngeal portion of the basicranium. In other specimens, such as *Pr. calchaquiorum* MACN-PV 5528, *Pro-macrauchenia* sp. MLP 29-X-10-16, and *S. bravardi* MACN-PV 13082, basicranial damage is extensive and it cannot be determined whether similar canals existed in these specimens, although the deep grooves emanating from the piriform fenestra are present. In her description, Fernández de Álvarez (1940) made mention of these apertures in *M. patachonica* MACN-PV 2 and interpreted them as channels for the auditory (= Eustachian) tubes, but this cannot be correct because they are situated well rostral to the middle ear and in any case pass into, not out of, the cranium (in her pl. 2, however, the arrow for "trompa de Eustaquio" is shown as emanating from damaged pneumatic cells in the pterygoid plate, which does not agree with her description).

Nor can canal X be the homolog of the alisphenoid canal of other mammals. The alisphenoid canal for the maxillary artery (or a branch derived therefrom) has an anterior and posterior opening, although the anterior opening may

merge with the sphenoorbital fissure or foramen rotundum and the caudal opening is frequently found in a common depression with the foramen ovale (e.g., Wible and Spaulding, 2013, and references herein cited). Moreover, a well-defined groove creasing the lateral aspect of the alisphenoid, in the form of a sulcus rather than a tube, was found in all members of the comparative set in which this area was represented, except for the *S. bravardi* MACN-PV 13082. If this channel did not conduct the maxillary artery per se, it may have transmitted a meningeal branch of the latter through foramen ovale. Apart from Macraucheniiidae, the groove in the alisphenoid is also seen in the Santacrucian protheriid *Tetramerorhinus cingulatum* MACN-A 5971. In this specimen there is also a relatively small aperture in the position of canal X.

What, then, is canal X and what is the relationship between it and the groove emanating from the piriform fenestra? We may begin with the identity of the latter. Although certainty is not possible, we infer from comparative considerations (MacPhee, 1981) that it probably housed the nerve of the pterygoid canal. This nerve, which transmits autonomic fibers related to the pterygopalatine ganglion (Hiatt and Gartner, 2001), usually runs within or along the pterygosphenoid suture to penetrate the rear wall of the pterygopalatine fossa, and it is always small in caliber. The unusual width of the groove in macraucheniiines suggests that it may have conducted something in addition. The essential basis for this assumption is as follows. Although canal X is differently positioned in IANIGLA-PV 29 and in *Pr. antiqua* MACN-PV 7986, in both specimens it is very large, much larger than would be necessary to transmit the nerve of the pterygoid canal alone. This can best be accounted for by assuming that the canal transmitted a hypertrophied vidian artery and perhaps also an associated vein. Other possibilities exist, such as an anastomotic branch from the ascending pharyngeal, but in view of the thinness of the evidence we prefer to simply name the inferred vessel the vidian or artery of the pterygoid canal, without

settling its homology. In the case of the latter specimen the apertures are also situated where one would expect the pterygoid canal to lie, i.e., at the junction of the central stem and dorsal sutural surface of the pterygoid.

Furthermore, because the fossa for the pterygopalatine ganglion is situated in close relation to the floor of the orbit, conditions in *Huayqueriana* would conform to those in other mammals if the nerve of the pterygoid canal simply ran across the opening of canal X to enter the pterygopalatine fossa. This does not, however, explain the comparatively large size of canal X nor its connection with the orbit. Intact vidian arteries in the adult stage are rare (or rarely identified) in mammals (MacPhee, 1981). Apparently even rarer is the situation in which the vidian forms an anastomotic link (arteria anastomotica) between the petrosal or cerebral portion of the internal carotid and the maxillary artery of the external carotid (De La Torre and Netsky, 1960). Typically, however, arteriae anastomoticae (which may arise in several different ways) do not possess their own foramina, but make their connections via foramen ovale. In artiodactyls such linkages, together with other associated retia, are thought to function as a countercurrent selective brain-cooling mechanism, with heat energy transferred from warm arterial blood to cooler venous blood (Caputa, 2004). Whether such a mechanism existed in *Huayqueriana* or other macraucheniiines is unknown and perhaps unknowable, as retia normally do not leave any diagnostic osseous indications of their existence (H. O'Brien, 2013, personal commun.). Phylogenetic bracketing does not resolve the issue, except in a negative sense, because cranial retia have not been identified in extant perissodactyls (Daniel et al., 1953; but see Du Boulay, 1999).

The jugular foramen (= foramen lacerum posterius; Fernández de Álvarez, 1940) is small and constricted. It opens rostroventrally, between the paracondylar process and the petrosal, and is hidden by the former in ventral view (fig. 16). There is a large piriform fenestra (= foramen lacerum anterius; Scott, 1910; Rusconi, 1932, Fernández de

Álvarez, 1940) (fig. 15), which intervenes between the anterior pole of the petrosal and the trailing edge of the alisphenoid (MacPhee, 1981). As in extant perissodactyls (e.g., *Equus*; Sisson and Grossman, 1975), the medial side of the petrosal is separated from the bones of the central stem by a broad fissure. This hiatus is continuous with the more medially situated piriform fenestra. *Huayqueriana* is similar, although the petrosal forms a tighter junction caudally with the basioccipital than in extant perissodactyls.

Although a separate opening for the internal carotid was not identifiable, the vessel may well have been present judging from comparative data on a variety of SANUs (Billet and Muizon, 2013; MacPhee, 2014; but for conditions in extant perissodactyls see Du Boulay, 1991). If present in the adult stage, it presumably passed into the endocranium through the piriform fenestra. However, no specific osteological features could be identified in CT scans that might support such an inference (MacPhee, 2014; Billet et al., 2015; see also Petrosal), although it is important to note that a certain amount of bone was lost during preparation in the relevant region.

The hypoglossal foramen (= condylar foramen of Scott, 1910; Rusconi, 1932; Fernández de Álvarez, 1940) is single and large, as in all macraucheniine specimens examined. It opens within a depression in front of the condyles, by which it is hidden in ventral aspect (fig. 15).

The occipital condyles are partially eroded and slightly dorsoventrally compressed in IANIGLA-PV 29. In lateral view (fig. 3A), the condyles do not reach the transverse level of the nuchal crest, differing in this regard from *X. bahiense*, *W. delacroixi*, and *M. patachonica*, whose condyles protrude beyond this level. In ventral view (fig. 3C), the condylar articular facets extend rostrally onto the basioccipital and are separated at the midline, similar to other macraucheniids (e.g., *Theosodon*, *Cr. normalis*, *S. bravardi*, *Pr. antiquua*, and *M. patachonica*; Scott, 1910; Rusconi, 1932; Dozo and Vera, 2010). The odontoid notch between the condyles is U-shaped, as in *Pr. calchaquiorum* (fig. 7C). This notch is narrower and V-shaped in *Pr.*

antiquua (fig. 7B), *S. bravardi* (fig. 9C), *W. delacroixi*, *X. bahiense*, and *M. patachonica* (Fernández de Álvarez, 1940). The foramen magnum is circular and directed caudoventrally.

The paracondylar process is broken on both sides of the skull. It is lateromedially wider than rostrocaudally long, and curves slightly rostrally. This process is more vertical and ventrally projecting in *Pr. antiquua* (fig. 5B), *S. bravardi* (fig. 9C), and *M. patachonica* (Fernández de Álvarez, 1940). In these species it protrudes below the plane of the basicranium.

In posterior view, the occipital is high, triangular, and ventrally broad (fig. 18). There is a vertical ridge descending from the nuchal crest, separating two lateral depressions that are laterally bordered by blunt occipital protuberances. The development of the occipital crest and associated protuberances varies among the specimens compared. In analogy to the horse, the bilateral depressions immediately under the nuchal crest and occipital protuberances should represent attachment points for the tendons of semispinalis capitis muscles (Sisson and Grossman, 1975).

The mastoid portion of the petrosal is wedged between the posttympanic and paracondylar processes, as in other litopterns (Scott, 1910; Rusconi, 1932; Soria, 2001). Between the mastoid and the paracondylar process there is a deep, almost vertical sulcus, the occipital groove, leading to the posttemporal canal, level with the dorsal border of the condyle (fig. 18). An opening in a similar position was described for *S. bravardi* (= mastoid foramen of Rusconi, 1932) and can be seen in the holotype of *H. cristata* as well as *O. zeballosi* MHIN-UNSL-GEO-V 465a, *Pr. antiquua*, and *M. patachonica* MACN-PV 2. As in other SANUs (MacPhee, 2014), the posttemporal canal, a primitive feature that transmits the arteria diploëtica magna in a diverse array of mammals (Wible, 1987), is confluent with the transverse sinus, which is mostly enclosed by bone and very large in *Huayqueriana* (figs. 11, 17A). In all probability the canal transmitted both an artery and a vein.

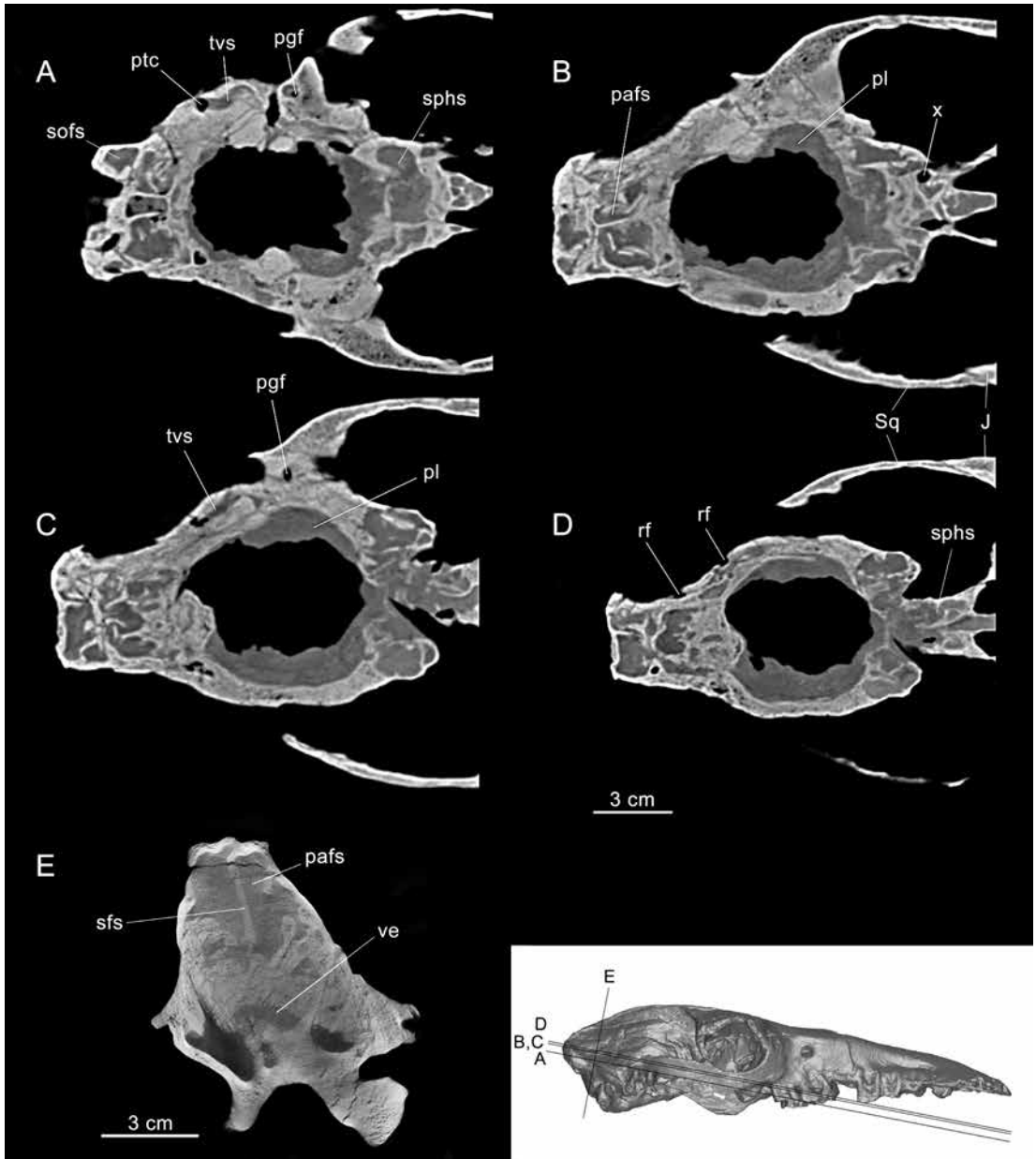


FIG. 17. A–E, Virtual subhorizontal and subcoronal sections of skull of *Huayqueriana* cf. *H. cristata* IANI-GLA-PV 29. Abbreviations: **J**, jugal; **pafs**, parietal extension of frontal sinus; **pgf**, postglenoid foramen; **pl**, position of piriform lobe; **ptc**, posttemporal canal; **rf**, foramina for rami temporales; **sfs**, septum sinuum frontalis; **sofs**, supraoccipital extension of frontal sinus; **sphs**, sphenoidal sinus; **Sq**, squamosal (zygomatic process); **tvs**, transverse sinus canal; **ve**, position of vermis. Letter x identifies canal X for nerve of pterygoid canal and vascular structures (see text).

It may be briefly noted that, although the posttemporal canal is located much more laterally and ventrally in macraucheniniines than in notoungulates (MacPhee, 2014), in the latter group it has the same relationship with the petrosal and occipital, and connects with the transverse sinus in the same way. Thus, we see no reason to question the homology of these openings in the different SANU orders.

IANIGLA-PV 29 exhibits an extra opening on the nuchal crest, probably a venous channel, at a more dorsal level on the occipital (fig. 18). Intracranially this canal is of small diameter and separate from the posttemporal canal. In dogs the named mastoid foramen transmits the occipital emissary vein, which drains the deep muscles of the neck into the sigmoid sinus (Evans, 1993), but the homology of this vessel in SANUs is problematic. As far as could be determined, this second aperture on the nuchal crest of IANIGLA-PV 29 is absent in the other macraucheniniine specimens examined.

MIDDLE EAR

The volume enclosed by the tympanic cavity is comparatively small in relation to skull size (fig. 19A). Its limits are difficult to reconstruct since much of the tympanic floor was evidently membranous and the area has been extensively prepared. Apart from the proximal hyoid (tympanohyal + stylohyal), the only element present for certain in the floor is the ectotympanic, which is preserved on both sides of the skull (fig. 15). The external acoustic meatus (fig. 15) is similar in width to that of the holotype of *H. cristata* (fig. 2C) and *Pr. calchaquiorum* (fig. 7C). In *Pr. antiquua* (fig. 7B) and *M. patachonica*, meatal width is somewhat greater.

ECTOTYMPANIC: As in *Rhinoceros* and *Tapirus* (van Kampen, 1905), as well as in other litopterns in which the ectotympanic is known (e.g., *Epitherium laternarium* [Soria, 2001]; *Theosodon* [Scott, 1910: pl. XVII]; *Huayqueriana cristata* holotype), this element in IANIGLA-PV 29 is only vaguely crescentic, lacking significant

medial and lateral flanges and forming neither an enclosing bulla nor a well-defined external acoustic meatus. In this regard litopterns are quite different from notoungulates (Patterson, 1932, 1934; Gabbert, 2004; MacPhee, 2014; Forasiepi et al., 2015b).

The rostral and caudal crura of the ectotympanic meet the alisphenoid in front and the mastoid/hyoid and paracondylar process behind, respectively, as in the holotype of *H. cristata*. In ventral view (fig. 15), the ectotympanic has three small, blunt protuberances, described further below. In front, a large process protrudes rostroventrally, as in the holotype of *H. cristata*, and exhibits a foramen at its base. Just caudal to this is another small process, broken on the left side, that points toward the meatus. The rearmost protuberance is medially oriented.

Related to these processes is the stump of the hyoid apparatus, which is closely appressed to the ectotympanic's caudal crus. Given its position it helps to delimit, together with the mastoid and paracondylar processes, an incomplete stylo-mastoid foramen for the exit of CN VII. This area is also the only place where the ectotympanic seems to have been attached to the skull (fig. 19A, left side). Whether there was also a rostral attachment to the anterior process of the malleus (= folian process), as occurs in *Rhinoceros* and *Tapirus* (van Kampen, 1905), cannot be determined. Although there must have been a tympanic membrane closing off the lateral aspect of the middle ear, its line of attachment on the ectotympanic is not obvious. The medial border of the left ectotympanic is slightly inflected for a short distance (fig. 19A, feature 2); this may represent the tympanic crest to which the membrane characteristically attaches in mammals.

On the same side there is another process (fig. 19A, feature 1) which extends in the form of a thin plate. The level of resolution in the illustrated segment is such that it cannot be determined whether this plate is ectotympanic or petrosal in origin, or something else entirely, such as an entotympanic. Although an entotympanic occurs in extant *Rhinoceros*, *Equus*, and possibly *Tapirus*

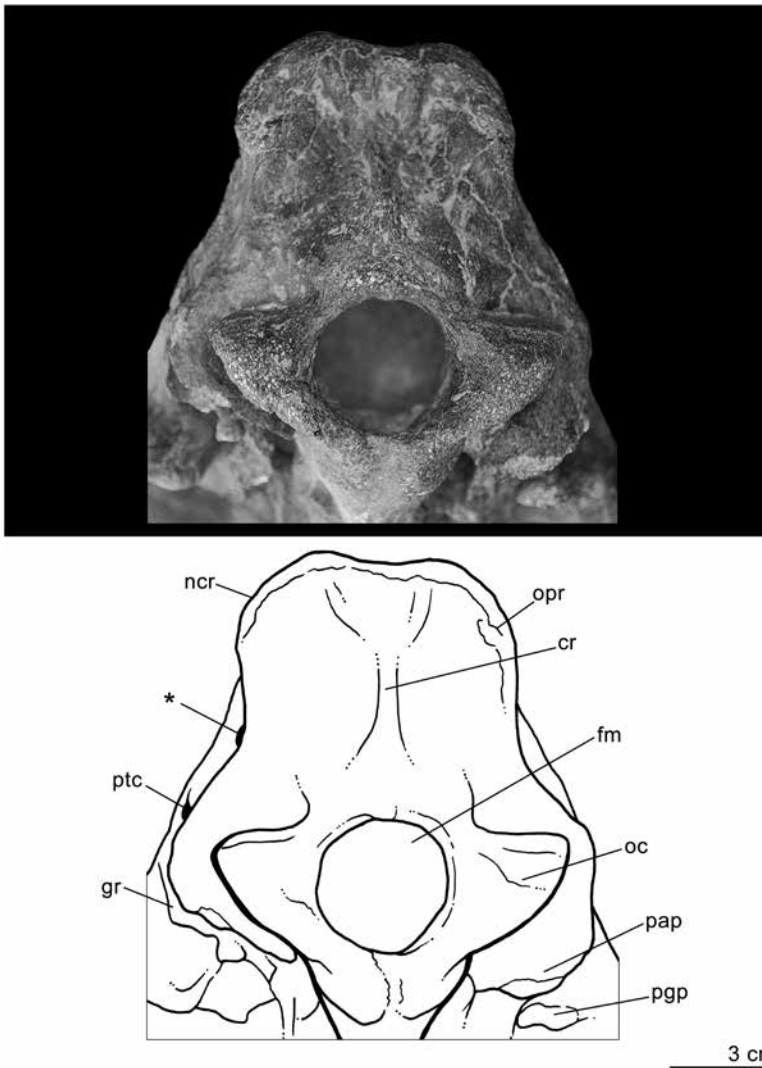


FIG. 18. Occipital view, skull of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **cr**, crest; **fm**, foramen magnum; **gr**, occipital groove; **ncr**, nuchal crest; **oc**, occipital condyle; **opr**, occipital protuberance; **pap**, paracondylar process of exoccipital; **pgp**, postglenoid process; **ptc**, posttemporal canal. Asterisk (*) identifies canal for vascular structures (see text).

(van Kampen, 1905; Maier et al., 2013), and apparently also in the notoungulate *Cochilius* (MacPhee, 2014), any conclusion concerning *Huayqueriana* will have to await better material.

In IANIGLA-PV 29, there is evidence that parts of some auditory ossicles were preserved. On the specimen's right side, a small piece of bone projects laterally from fenestra vestibuli

(fig. 19B). It connects with a second piece, the dorsal end of which is lodged in a small depression on the tympanic roof medial to the crista parotica. Resolution is poor, but it seems plausible that the features seen on the scans include one of the stapedial crura and the incudo-stapedial joint. This is the only known example of ossicle preservation in a litoptern.

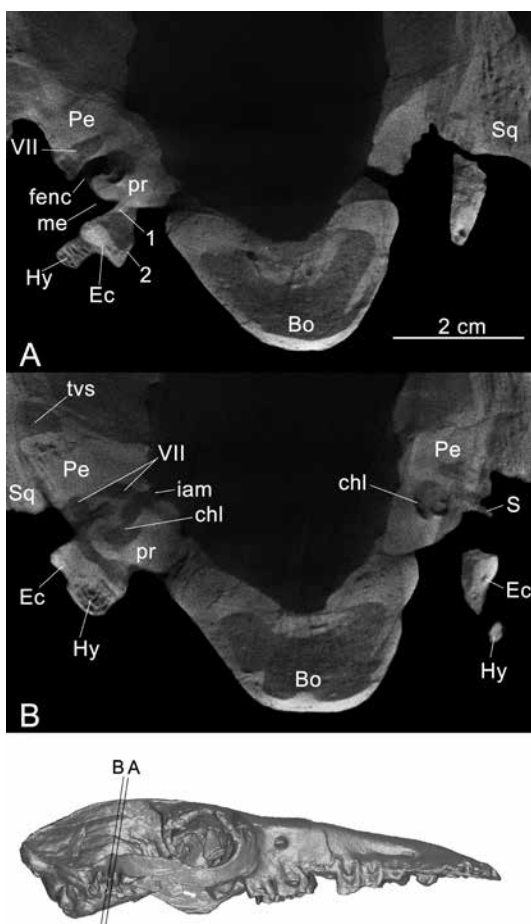


FIG. 19. **A–B**, Virtual subcoronal sections of basicranium of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Abbreviations: **Bo**, basioccipital; **chl**, cochlea; **Ec**, ectotympanic; **fenc**, fossula fenestrae cochleae; **Hy**, hyoid (tympanohyal + stylohyal); **iam**, internal acoustic meatus; **me**, middle ear space; **Pe**, petrosal; **pr**, promontorium; **S**, stapes; **Sq**, squamosal; **tvs**, transverse sinus; **VII**, passageway of the facial nerve (CN VII); **1**, RTPP contacting ectotympanic; **2**, crista tympanica?

PETROSAL: As the density contrast in CT segments was low, we were unable to produce high-quality virtual 3D reconstructions of the petrosal. However, the tympanic, oblique, and cerebellar views of the reconstructed right petrosal presented in figure 20 are adequate for illustrating and describing principal anatomical features (fig.

20A is provided with an orientation guide). In order to adequately visualize deeply recessed features in tympanic or ventral aspect, it was necessary to virtually cut away much of the crista parotica, pars canalicularis, and tympanic floor (tympanohyal, ectotympanic). However, most of these features can be seen in section in the segmental views (fig. 19). The aperture in the lateral sidewall of the promontorium (asterisk in fig. 20A, B) is a preparation artifact, not a preexisting structure. It opens into the cochlear canal.

The rounded promontorium, housing the cochlea (e.g., MacIntyre, 1972; Meng and Fox, 1995), dominates the tympanic aspect (fig. 20A). The lateral and caudal surfaces of the promontorium are interrupted by the fenestra vestibuli and the fossula fenestrae cochleae, respectively. The fenestra cochleae, which is situated within the fossula and is not externally visible from this angle (MacPhee, 1981), internally supports the secondary tympanic membrane. The fenestra cochleae is separated from the fenestra vestibuli by a thick, semivertical crista interfenestralis. The fenestra vestibuli, which holds the footplate of the stapes in life, is similar in area to the fenestra cochleae. The stapedial ratio (~ 1.8) is larger than that of other litopterns that have been examined, including *Macrauchenia* (1.5; Billet et al., 2015).

The caudal end of the tympanic cavity is capacious, but no features associated with it (e.g., stapedius muscle fossa, fossa incudis) can be securely identified on the reconstruction. Because the line of the cut through the mastoid area passes through this region, it is not evident that it is, in fact, framed by the petrosal (in the form of the caudal tympanic process of the petrosal, CTPP).

Rostromedially, there is a very low, discontinuous crest on the ventral surface of the promontorium that continues onto its medial side. Inspection of relevant segments indicates that the arc of curvature of the crest corresponds to that of the medial margin of the ectotympanic, and that the two were in close contact during life. This feature may therefore be identified as a rather insignificant rostral tympanic process of

the petrosal (RTPP), the ventral edge of which helped to form the tympanic/petrosal suture. A somewhat more developed crest is seen in a similar position in Paleogene litoptern fossils referred to *Miguelsoria* (Billet et al., 2015). The RTPP has been identified in a wide variety of mammals (e.g., MacPhee, 1981; Wible 2003, 2008, 2012), although the derived eutherian condition, in which the process forms most or all of a well-developed bulla, is largely limited to primates and possibly some of their close relatives.

Medial to the RTPP, on the medial edge of the petrosal, there is shallow groove that may have held the inferior petrosal sinus (SIPS). This structure is very common in mammals, as the sinus is the major connector between pharyngeal venous plexuses, the cavernous sinus, and the internal jugular vein on the ventral surface of the skull. Salient crests bordering a similar sulcus have been described for *Miguelsoria* (Billet et al., 2015).

The area immediately lateral to the preparation artifact in IANIGLA-PV 29 is also damaged (fig. 20B), and in order to visualize structures it was necessary to digitally open up the area related to the cavum supracochleare (for the geniculate ganglion and first part of the facial nerve). Two sulci emerge from the cavum supracochleare, one running caudomedially to the incomplete stylomastoid foramen (sulcus for the facial nerve), and the other, deeper groove running rostromedially to the hiatus Fallopii (hiatus for the greater petrosal nerve). The foramen faciale (FF), the aperture in the tympanic roof by means of which the second or tympanic part of the facial nerve enters the true tympanic cavity from the cavum supracochleare, is represented in the reconstruction by a notch corresponding to the foramen's dorsal border (fig. 20B).

The cerebellar view of the petrosal (fig. 20C) is dominated by two main structures: the internal acoustic meatus (IAM) rostrally and the subarcuate fossa (SAF) in the caudal part of the bone. The deep meatus is subdivided as usual into the superior acoustic foramen (FAS) laterally and the inferior acoustic foramen (FAI)

medially (fig. 20C), the two being separated by a thick crista transversa. In *Miguelsoria* (Billet et al., 2015) the IAM appears to have been shallower. The FAS splits into two openings: a large rostral aperture, leading to the cavum supracochleare, for the facial nerve (CN VII); and a smaller caudal opening corresponding to the channel for a branch of the vestibular nerve (CN VIII), which innervates the membranous labyrinth (e.g., Wible, 2003). The FAI also subdivides into a large rostral aperture for bundles of the vestibular nerve and a smaller caudal area perforated by many small foramina (cribriform tract, for fascicles of the cochlear nerve). The caudal border of the internal acoustic meatus bears a tiny aperture that presumably corresponds to the foramen singulare for passage of some fibers of the vestibular nerve (CN VIII) (Meng and Fox, 1995).

The subarcuate fossa, as seen in the 3D model of the petrosal (fig. 20C), is even shallower than in the material referred to *Miguelsoria* (Billet et al., 2015). This is in agreement with the small size of the cast of the cerebellar paraflocculus, the structure that it would have housed in life (e.g., MacIntyre, 1972) (see Endocast).

The external opening of the cochlear canaliculus (= aqueductus cochleae) for the perilymphatic duct (see Osseous Inner Ear) is small and located on the medial side of the petrosal, in relation to the notch marking the border of the jugular foramen. A single cochlear canaliculus is present in IANIGLA-PV 29, differing from some specimens referred to *Miguelsoria* in which there is a small accessory aperture (probably vascular) accompanying the cochlear canaliculus (Billet et al., 2015).

OSSEOUS INNER EAR

The endocast of the left osseous labyrinth of IANIGLA-PV 29 is illustrated in figure 21. As in all mammals, the labyrinth consists of a series of interconnected spaces within the petrosal, including the cochlea, vestibule, and the three semicircular canals (e.g., MacIntyre, 1972; Ekdale, 2013, 2016). The cochlear canal

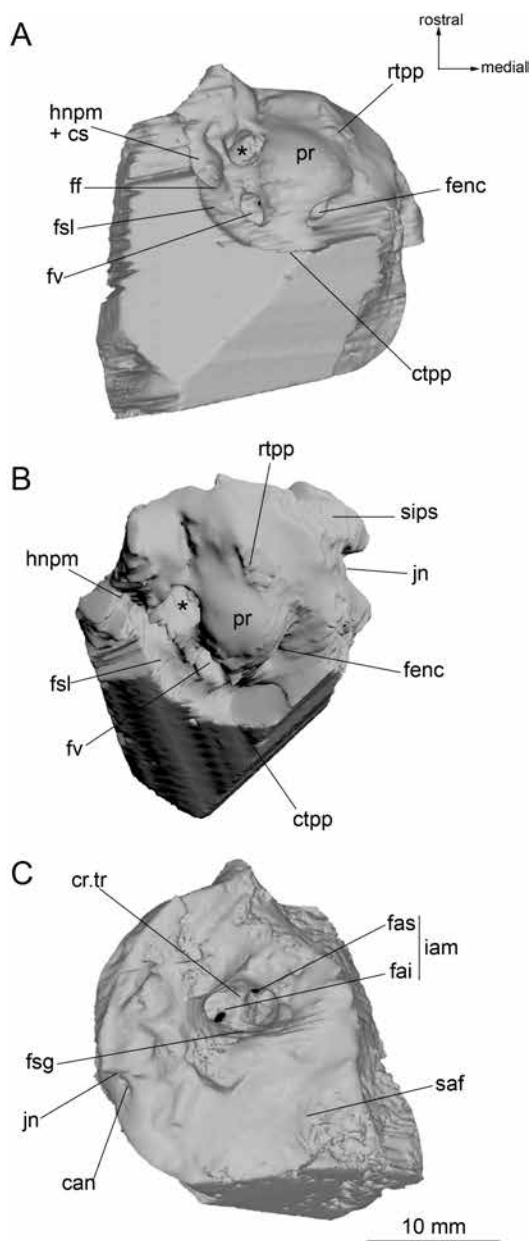


FIG. 20. Model of right petrosal of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, based on 3D reconstructions from micro-CT data, in **A**, tympanic, **B**, oblique, and **C**, cerebellar views. In order to adequately expose the facial sulcus (**fsl**) and track of the greater petrosal nerve (**hnpm**), it was necessary to digitally remove the bone obscuring structures, including the sidewall of the cavum supracochleare (**cs**) (see text). Other abbreviations: **can**, external aperture of cochlear canaliculus; **cr.tr**, crista transversa; **ctpp**, caudal tympanic process of petrosal; **fai**, inferior acoustic foramen; **fas**, superior acoustic foramen; **fenc**, fossula fenestrae cochleae; **ff**, foramen faciale; **fsg**, foramen singulare; **fv**, fenestra vestibuli; **iam**, internal acoustic meatus; **jn**, notch for jugular foramen; **pr**, promontorium; **rtp**, rostral tympanic process of petrosal; **saf**, subarcuate fossa; **sips**, sulcus for inferior petrosal sinus. Asterisk (*) indicates preparation artifact.

forms the largest part of the osseous labyrinth by volume (table 3). The canal is coiled in *Huayqueriana*, as is universally the case in therians (Luo et al., 2010); it expresses a total helical rotation of 795° , equivalent to 2.2 full turns (fig. 21F; table 3). All litopterns measured to date exhibit similar values, including Early Eocene taxa (2.25 to 2.4 turns), “*Proterotherium*”² (2.1), *Diadiaphorus* (2.25), and *Macrauchenia* (2.3) as measured by Billet et al. (2015). The notoungulates *Notostylops* (2.25; Macrini et al., 2010) and *Hegetotherium* (2.5; Simpson, 1936; Macrini et al., 2010) are also similar, whereas in some other notoungulates (*Altitypotherium*, *Pachyrhinos*, and *Cochilius*) the degree of coiling is less, ~ 2 turns (Macrini et al., 2013).

The first half of the basal cochlear turn is not tightly appressed to the rest of the coil, which is a similarity to conditions previously described for other litopterns (Billet et al., 2015). By contrast, succeeding turns are tightly packed. The primary and secondary osseous spiral laminae, the presence of which is a plesiomorphic therian trait (Meng and Fox, 1995), incompletely separate the cochlear canal into two parts, the scala tympani and scala vestibuli. In IANIGLA-PV 29 the primary lamina is represented by a feeble groove on the inner wall of the cochlea, while the deeper secondary lamina on the outer surface of the spiral can be seen along the basal turn to a point $\sim 510^\circ$ away from the fenestra cochleae (i.e., 0°) (fig. 21E), similar to other litopterns (Billet et al., 2015). In profile, the cochlea of IANIGLA-PV 29 has a height that is about two-thirds of its width (table 3).

² Identified as *Proterotherium* by Billet et al. (2015), but see taxonomy in Soria (2001). In view of the taxonomic uncertainty surrounding the allocation of the specimens studied, it is here referred to as “*Proterotherium*.”

TABLE 3

Linear and angular measurements of the digital inner ear of *Huayqueriana* cf. *H. cristata* (IANIGLA-PV 29)

Abbreviations: L, length; W, width.

Number of cochlear turns ¹	2.2
Cochlea height	3.93mm
Cochlea width	5.77mm
ASC L/ ASC W ²	7.84mm / 7.87mm
ASC L/W	0.99
LSC L / LSC W ²	7.09mm / 6.97mm
LSC L/W	1.02
PSC L / PSC W ²	7.68mm / 7.71mm
PSC L/W	0.99
Angle between ASC and PSC ³	88°
Angle between the PSC and LSC ⁴	90°
ASC dorsal projection/PSC height ratio ⁵	0.128

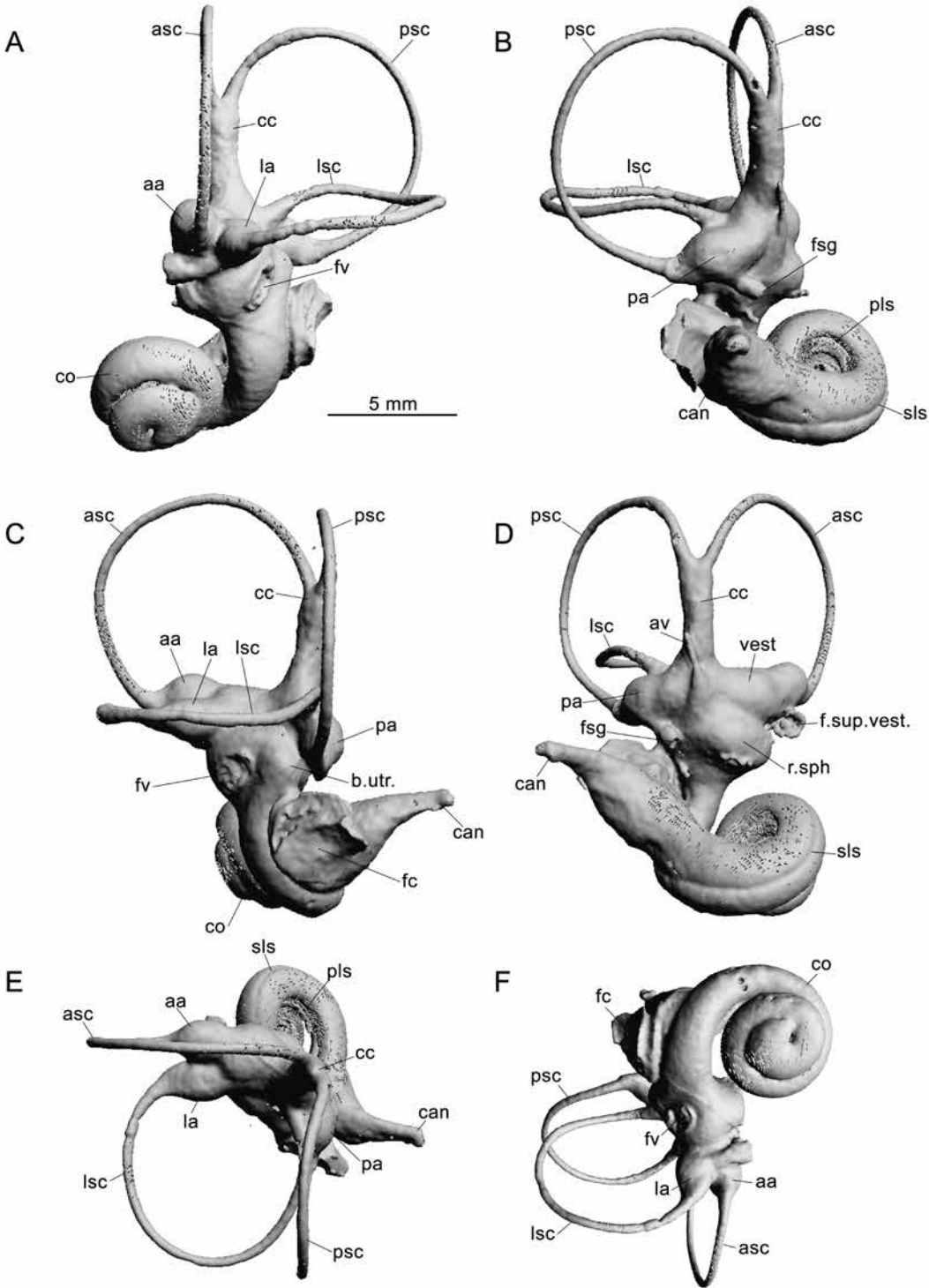
¹ Counted following the protocol of West (1985).
² Measurements taken from the inner side of the SC following the protocol of Schmelzle et al. (2007).
³ Measured on a plane parallel to the LSC. The origin of the angle is at the crus commune (Schmelzle et al., 2007).
⁴ Measured on a plane parallel to the ASC (Schmelzle et al., 2007).
⁵ Distance between the dorsal-most point of the ASC to the dorsal-most point of the PSC/ height of PSC measured in straight line from the LSC to the dorsal-most point of the PSC (Schmelzle et al., 2007).

The main axis of the fenestra vestibuli is oriented caudodorsally-rostroventrally, and is positioned ventral and slightly posterior to the lateral ampulla (fig. 21A, C). The external aperture of the fossula fenestrae cochleae is a section of a distorted cone (rather than oval, as usually stated) and larger than that of the fenestra vestibuli, as is also the case in other described litopterns and the notoungulate *Notostylops* (Macrini et al., 2013; Billet et al., 2015).

The cochlear canaliculus (= aqueductus cochleae), for the membranous perilymphatic duct, opens within the endocranium caudomedial to the fossula fenestrae cochleae. It projects beyond the PSC as seen in dorsal view (fig. 21E), as in other described litopterns and *Notostylops* (Macrini et al., 2010; Billet et al., 2015), in contrast to *Pachyrukhos* in which it does not extend as far (Macrini et al., 2013). However, it should be noted that the difference may be size related, in the sense that *Pachyrukhos*, a very small

notoungulate, does not exhibit greatly thickened capsular walls. In consequence, the perilymphatic duct has a shorter journey to the endocranium, reflected in the abbreviated length of its osseous canal.

The vestibule contains the membranous utricle and saccule (Meng and Fox, 1995; Macrini et al., 2010; Ekdale, 2016). In IANIGLA-PV 29 the vestibule is rostrocaudally elongated, parallel to the main axis of the LSC. The recessus sphericus of the vestibule, which contains the saccule in life, forms a significant bulge in the vicinity of the fenestra vestibuli in posteromedial view (fig. 21D). There is a complementary bulge on the utricular side of the vestibule, but it is smaller when seen in lateral view (fig. 21C). In posterior view (fig. 21B, D) the canal for nervus ampullaris posterior (CN VIII) runs ventrally from the posterior ampulla to communicate with foramen singulare. The foramen for the superior vestibular area, which conducts the superior part of the



vestibular (utrículoampullary) nerve to the utricular macula, is situated as usual just beneath the ampulla of the anterior semicircular canal (fig. 21D). The narrow aqueductus vestibuli, for the membranous endolymphatic duct, projects dorsally from the vestibule, parallel to crus commune (fig. 21D).

The three semicircular canals are located in the pars canalicularis of the petrosal. The ASC forms part of the rim of the subarcuate fossa, the PSC contributing to its medial border. As in other described litopterns (Billet et al., 2015), the ASC, PSC, and LSC of IANIGLA-PV 29 are notably gracile, much thinner than the crus commune. The ASC and PSC are uniplanar while the LSC is slightly undulating, curving dorsally and then ventrally before connecting to the posterior ampulla (fig. 21B). The semicircular canals are arranged approximately at mutual right angles (table 3). The shape, relative size, and radius of curvature of the semicircular canals vary among litopterns and notoungulates (see Macrini et al., 2010, 2013; Billet et al., 2015).

The arc of the ASC projects slightly above that of the dorsalmost point on the PSC (fig. 21A; table 3), similar to the notoungulate *Altitypothierium* (Macrini et al., 2013). The degree of projection is apparently greater in *Macrauchenia*, proterotheriids (e.g., “*Proterotherium*” and *Diadiaphorus*; cf. Billet et al., 2015: fig. 17), and the notoungulates *Pachyrhinos* and *Cochilius* (Macrini et al., 2013). In dorsal view (fig. 21E), the arc of the LSC is almost level with the lateralmost projection of the PSC, similar to *Diadiaphorus* and the notoungulates *Notostylops*, *Altitypothierium*, and *Pachyrhinos*, but in contrast to the more extended PSC of “*Proterotherium*,” *Macrauchenia*, and the notoungulate *Cochilius* (Macrini et al., 2010, 2013; Billet et al., 2015).

The three ampullae lie in close proximity to the junction of the semicircular canals with the ventricle. The anterior ampulla is on the rostro-ventral arm of the ASC and dorsolateral to the vestibule; the lateral ampulla is on the anterior arm of the LSC and dorsolateral to the vestibule, and the posterior ampulla is on the ventral arm of the PSC and ventromedial to the vestibule. There is no secondary crus commune (fig. 21B–D), or the short canal formed by the union of the posterior arm of the LSC and the ventral arm of the PSC (Schmelzle et al., 2007). The secondary crus commune is also absent in *Macrauchenia* and *Diadiaphorus*, and the notoungulates *Notostylops*, *Altitypothierium*, *Pachyrhinos*, and *Cochilius*, but it is present in Early Eocene litopterns (Macrini et al., 2010, 2013; Billet et al., 2015). The secondary crus commune is a plesiomorphic trait for mammals (e.g., Meng and Fox, 1995; Ekdale, 2016), evidently retained in basal litopterns but subsequently lost in later members of the group (Billet et al., 2015).

The crus commune is formed by the posterior arm of the ASC and the anterior arm of the PSC, as in all mammals (Ekdale, 2013). The crus commune is shorter than the length of the two individual SCs, as in other litopterns (excepting some specimens of *Miguelsoria* in which the ASC but not the PSC is longer; Billet et al., 2015). Among notoungulates the described variation seems to be greater than in litopterns (Macrini et al., 2010, 2013).

CRANIAL PNEUMATIZATION

In contrast to the relatively small volume of the middle ear, in at least some macraucheniids pneumatization of the cranial vault and basicra-

FIG. 21. Model of osseous labyrinth endocast of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, based on 3D reconstructions from micro-CT data, in **A**, anterior, **B**, posterior, **C**, lateral, **D**, posteromedial, **E**, dorsal, and **F**, ventral views. Abbreviations: **aa**, anterior ampulla; **asc**, anterior semicircular canal; **av**, aqueductus vestibuli; **b.utr.**, bulge for utricle; **can**, cochlear canaliculus; **cc**, crus commune; **co**, cochlea; **f.sup.vest.**, foramen for superior vestibular area; **fc**, fossula fenestrae cochleae; **fsg**, foramen singulare; **fv**, fenestra vestibuli; **la**, lateral ampulla; **lsc**, lateral semicircular canal; **pa**, posterior ampulla; **pls**, primary bony lamina sulcus; **psc**, posterior semicircular canal; **r.sph.**, recessus sphericus of vestibule (for saccule); **sls**, secondary bony lamina sulcus; **vest**, vestibule

nium is very marked indeed (figs. 10, 11, 17), as originally noted by Rusconi (1932) for *S. bravardi* MACN-PV 13082. Cranial pneumatization may have reached an ordinal maximum in *M. patachonica*. In *M. patachonica* MACN-PV 2, the tables of the cranial vault, central stem, pterygoid plates, and lateral aspect of the alisphenoid are all massively inflated (fig. 11). In IANIGLA-PV 29, pneumatization of the vault likewise affects the entire dorsum of the skull, from the frontals to the supraoccipital, although on a smaller scale (figs. 10, 11). Segmental evidence indicates that remodeling activity in the vault proceeded during ontogeny from the paranasal frontal sinuses, which progressively invaded the parietals, squamosals, and supraoccipital. In *Huayqueriana* (figs. 10, 11), the enormous right and left frontal sinuses are separated for their entire length by a septum (septum sinuum frontalem; fig. 17E), as in *Equus* (Sisson and Grossman, 1975) and probably many other perissodactyls (e.g., tapir; Witmer et al. 1999: fig. 7). Although scan resolution is too low for detailed interpretation, pneumatic activity within the basicranium of IANIGLA-PV 29 appears to have spread from the nasopharynx into the bones comprising the central stem, probably via outpocketings from the auditory tube separate from those inflating the middle ear (cf. pneumatization in some other SANU taxa; MacPhee, 2014).

As already mentioned, middle ear pneumatization in *Huayqueriana* and other Litopterna is not extensive (fig. 19A, B). Although it is frequently stated that litopterns lack an expanded epitympanic sinus (Roth, 1903), this is not the case for all taxa. Rusconi (1932) identified an epitympanic sinus caudodorsal to the external acoustic meatus in *S. bravardi*. Micro-CT data reveal that *Tetramerorhinus* also exhibits a small swelling above the tympanic cavity (S. Hernandez del Pino, pers. obs.). The difference is better understood as one of scale, with the epitympanic sinuses of litopterns usually being inapparent externally (unlike smaller notoungulates such as mesotheriids and pachyrukhines; cf. Patterson, 1934; MacPhee, 2014).

CEREBRAL ENDOCAST

To enhance accuracy in description, where useful we differentiate between actual brain structures (e.g., olfactory bulbs) as they would appear in vivo and the corresponding, but obviously nonisomorphic, portions of the endocast by adding a subscript “c” to the structure’s name (e.g., olfactory bulbs_c). This distinction seems unnecessary to make for nonneural tissues such as blood vessels.

The only previously published endocast of any litoptern is that of *Tetramerorhinus lucarius* (= *Proterotherium cavum*) AMNH 9245, described by Simpson (1933) (fig. 23; appendix 1; taxonomy follows Soria, 2001, p. 42). Using newly acquired segmental data, we extend and reevaluate his notes on this specimen in light of complementary micro-CT evidence for IANIGLA-PV 29 (fig. 22).

ENDOCAST FLEXURE (EF): EF is defined as the angle subtended by a line running from the center of the olfactory bulb_c to the center of the hypophysis_c, and another running from the latter point to the center of the foramen magnum (Macrini et al., 2006). In IANIGLA-PV 29 the EF value is 33° (table 4). The width/length ratio of the endocast (excluding the medulla_c) is 0.69, and the height/width ratio is 1.13 (table 4).

FOREBRAIN REGION: In life, the most anterior structures of the endocast would have housed the olfactory bulbs and associated meninges (Macrini et al., 2007b). The bulbar portion of the endocast is small in IANIGLA-PV 29, contributing only ~4% of the total volume (table 4). Bulbar contribution in *Te. lucarius* is also relatively small, although larger than previously thought (Simpson, 1933; fig. 23). In IANIGLA-PV 29 each bulb_c is subspherical (left side: width/length ratio, 1.45; height/length ratio 1.6), aligned with the main axis of the ipsilateral cerebral hemisphere_c, and widely separated from its partner at the midline. The circular fissure (= fissure endorhinalis of Simpson, 1933) separating the olfactory bulbs_c from the cerebral hemispheres_c is narrow and deep. The endocast lacks any indication of either accessory olfactory bulbs or olfac-

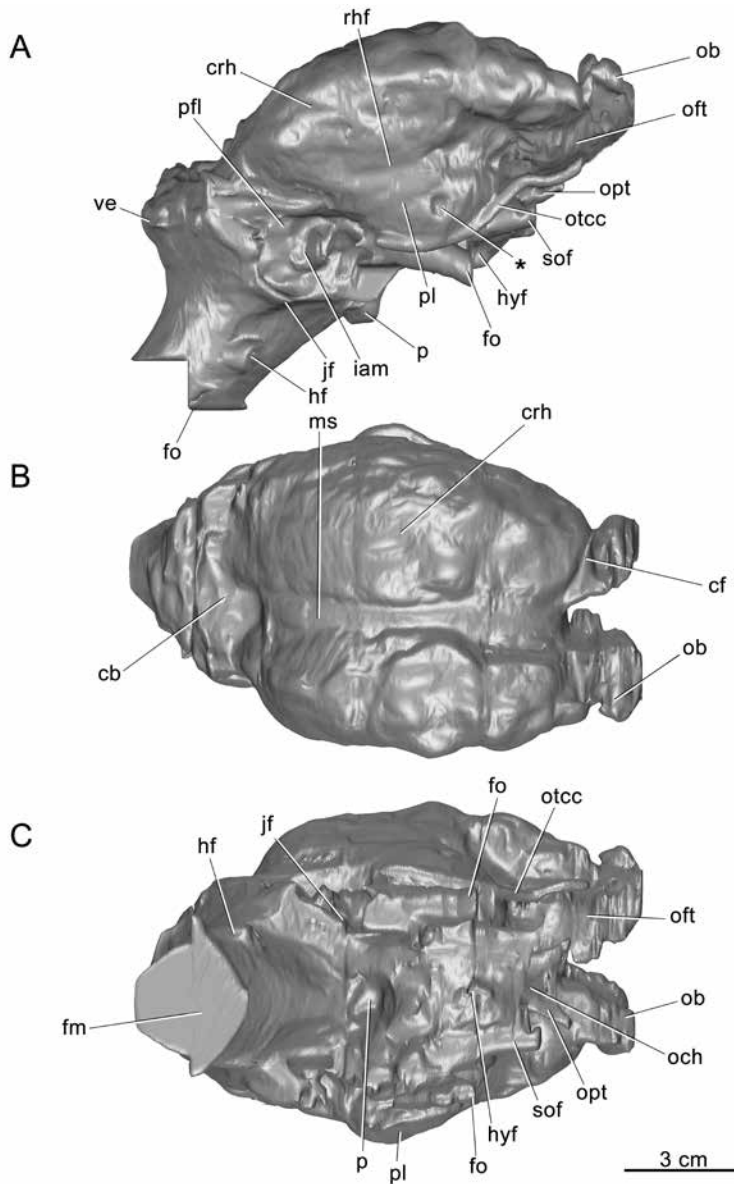


FIG. 22. Cerebral endocast of *Huayqueriana cf. H. cristata* IANIGLA-PV 29, in **A**, left lateral, **B**, dorsal, and **C**, ventral views. Abbreviations: **cb**, cast of cerebellum; **cf**, circular fissure; **crh**, cast of cerebral hemisphere; **fm**, spinal cord exiting foramen magnum; **fo**, mandibular division of trigeminal nerve (CN V₃) transiting foramen ovale; **hf**, hypoglossal nerve (CN XII) transiting hypoglossal foramen; **hyf**, cast of hypophyseal fossa; **iam**, facial and vestibulocochlear nerves (CN VII and CN VIII) transiting internal acoustic meatus; **jf**, glossopharyngeal, vagus, and accessory nerves (CN IX, CN X, and CN XI) transiting jugular foramen; **ms**, median sulcus; **ob**, cast of olfactory bulb; **och**, optic chiasm; **oft**, olfactory tract; **opt**, optic tract (CN II); **otcc**, cast of orbitotemporal canal; **p**, pons; **pfl**, cast of paraflocculus; **pl**, cast of piriform lobe; **rhf**, rhinal fissure; **sof**, oculomotor, trochlear, ophthalmic and maxillary divisions of trigeminal, and abducens nerves (CN III, CN IV, CN V_{1,2}, CN VI) leading to sphenoorbital fissure; **ve**, cast of vermis. Asterisk (*) indicates depression on right lateral piriform lobe.

TABLE 4
Digital endocast of *Huayqueriana* cf. *H. cristata* (IANIGLA-PV 29):
Linear, angular, and volumetric measurements¹

General		
Endocast, volume ²		217290 mm ³
Endocast, flexure		33°
Endocast, maximum length	128 mm (oblique) / 119 mm (straight)	
Endocast, maximum width		82.3 mm
Endocast, maximum height		73 mm
Bilateral structures		
Olfactory bulbs, R + L volume		8332 mm ³
Olfactory bulbs, R + L maximum width		48.4 mm
Olfactory bulb, R anteroposterior length		12 mm
Olfactory bulb, L anteroposterior length		15 mm
Olfactory bulb, R maximum height		23mm
Olfactory bulb, L maximum height		19 mm
Piriform lobes, R + L volume		19738 mm ³
Cerebral hemispheres, R + L volume		124276 mm ³
Cerebral hemisphere, R anteroposterior length		79.2 mm
Cerebral hemisphere, R maximum width		35.3 mm
Cerebral hemisphere, R maximum height		32.6 mm
Cerebral hemisphere, L anteroposterior length		81 mm
Cerebral hemisphere, L maximum width		42 mm
Cerebral hemisphere, L maximum height		36 mm
Midline structures		
Hypophysis, volume		5228 mm ³
Hypophysis, anteroposterior length		18.5 mm
Hypophysis, maximum width		16 mm
Hypophysis, maximum height		6 mm
Cerebellum, volume		48404 mm ³

¹As computed by the program 3DSlicer. ²Including olfactory bulbs but excluding medulla.

tory tracts projecting from the cortex (Macrini et al., 2007a, 2007b).

The cerebral hemispheres_c represent 57% of the total endocast volume (table 4). They are separated sagittally by a wide median sulcus. As already mentioned, the cerebral hemispheres are asymmetrical due to taphonomic deformation. The less affected right cerebral hemisphere_c is ovoid with a width/length ratio of 0.44 and a

height/length ratio of 0.41 (table 4). The cortical surface is slightly gyrencephalic, with very shallow sulci or gyri. Conspicuously lacking are the well-marked and comparatively deep subparallel dorsal sulci seen on the endocast of *Te. lucarius* (Simpson, 1933; fig. 23B). In contrast, the rhinal fissure is well marked in both endocasts.

The piriform lobes_c are laterally protruding, as in *Te. lucarius* (Simpson, 1933), and represent

9% of total endocast volume. In Simpson's (1933: fig. 3) illustration of *Te. lucarius*, there are two roughly circular depressions on each lobe_c, one ventrolateral and the other ventral. The ventral depression is absent in the 3D model of *Te. lucarius* (fig. 23C), suggesting that this feature in Simpson's plaster version was a casting artifact. In *Huayqueriana*, there is only a single circular depression (?preservation artifact), situated on the right piriform lobe (fig. 22A) and similar in position to one on the new endocast of *Te. lucarius* (fig. 23A).

The hypophyseal region_c, the space occupied in vivo by a portion of the pituitary gland (Macrini et al., 2007a), is represented by a small, shallow bulge, about 2.5 % of total endocast volume. It is situated in the middle of the ventral floor, level with the anterior border of the piriform lobe_c and slightly anterior to foramen ovale (fig. 15). The hypophysis_c exhibits a medial keel, but this is probably an artifact of preservation due to damage affecting the floor of the hypophyseal fossa. There is no bony indication of the passage of the cerebral carotid artery, such as the sulcus sometimes found in mammals in close relation to the hypophyseal fossa (MacPhee, 1981).

HINDBRAIN REGION: The depression that separates the cerebrum_c and cerebellum_c is notably shallower in IANIGLA-PV 29 (fig. 22B) than in *Te. lucarius* (Simpson, 1933), which is consistent with a negligible or absent tentorium cerebelli (Macrini et al., 2007b). The volume of the cerebellum_c forms 22% of the total endocast, which is smaller than its contribution in *Te. lucarius* (Simpson, 1933). Interestingly, in IANIGLA-PV 29 the cerebellum_c is narrower than any part of the cerebrum_c, whereas in dorsal view in *Te. lucarius* the former is as wide as the anterior portion of the cerebral hemisphere_c (Simpson, 1933).

In IANIGLA-PV 29 it is difficult to separate the impression for the vermis_c from that for the cerebellar hemispheres_c, but the scan indicates that in dorsal view the vermis_c is roughly semi-spherical (fig. 17E). The surfaces of the cerebellar hemispheres_c are smooth. Meninges, venous sinuses, and other structures (e.g., arachnoid

granulations) affect endocast topography and consequently there is no direct correlation between the apparent degree of lissencephaly seen on hemisphere casts and actual, in vivo surfaces (Macrini et al., 2007a). The paraflocculus_c, caudodorsal to the internal acoustic meatus on the lateral face of the cerebellar hemisphere_c, is less protruding than in *Te. lucarius* (Simpson, 1933). This correlates with the notable shallowness of the subarcuate fossa (see Petrosal).

The deepest endocast structure distinguishable in ventral view is the pons_c, which projects from the ventral surface as an oval bulge. Posteriorly, the medulla oblongata_c, circular in cross section, exits the skull via the foramen magnum.

NERVES AND VESSELS: Conspicuous nerves and blood vessels on the endocast of IANIGLA-PV 29 may be briefly noted.

Anterior cranial fossa. The short optic chiasm_c, formed by the grooves that carried the right and left optic nerves_c (CN II), is clearly evident on the border between the anterior and middle cranial fossae, although somewhat deformed. The optic nerves_c are relatively small in diameter and diverge at an angle of ~88° (fig. 22C).

Middle cranial fossa. The sphenoorbital fissure (= anterior lacerate foramen of Simpson, 1933) is the aperture through which the oculomotor (CN III), trochlear (CN IV), ophthalmic (CN V₁), maxillary (CN V₂), and abducens (CN VI) nerves typically leave the skull in eutherians (MacPhee, 1994). Because of their size and lateral disposition, the divisions of CN V are usually the only cranial nerves that can be separately identified in the middle cranial fossa, and this is the case in IANIGLA-PV 29. The casts of the tracks of CN V₁ and CN V₂ converge at the sphenoorbital fissure, while that for the mandibular nerve (CN V₃) passes through a distinct foramen ovale, as in other SANUs (Gabbert, 2004; Forasiepi et al., 2015b). As noted in the description of the skull, IANIGLA-PV 29 lacks a foramen rotundum providing an independent exit for the maxillary nerve (CN V₂). Scott (1910: 18) suggested that, in *Diadiaphorus* and other litopterns, CN V₂ and CN V₃ exited together through

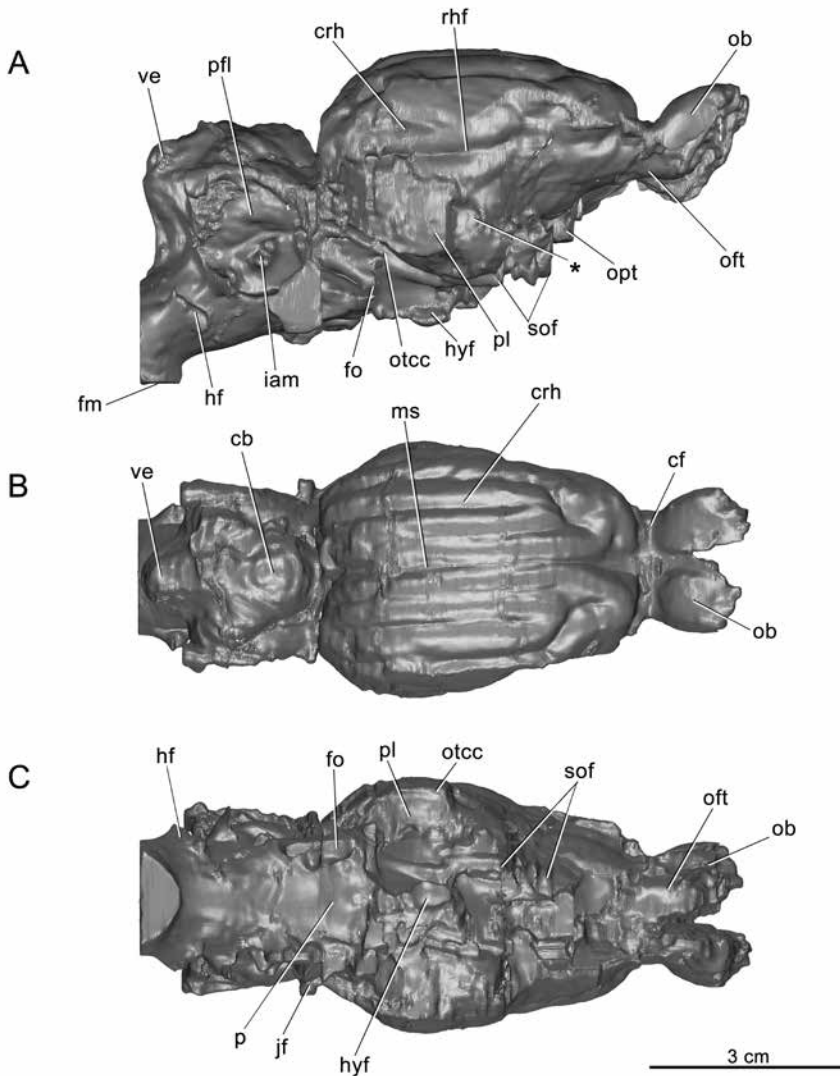


FIG. 23. Cerebral endocast of *Tetramerorhinus lucarius* (= *Proterotherium cavum* AMNH 9245) from Santa Cruz Formation, Santa Cruz, in **A**, left lateral, **B**, dorsal, and **C**, ventral views. Abbreviations: **cb**, cerebellum; **cf**, circular fissure; **crh**, cerebral hemisphere; **fm**, spinal cord exiting foramen magnum; **fo**, mandibular division of trigeminal nerve (CN V₃) exiting foramen ovale; **hf**, hypoglossal nerve (CN XII) exiting hypoglossal foramen; **hyf**, hypophyseal fossa; **iam**, facial and vestibulocochlear nerves (CN VII and CN VIII) transiting internal acoustic meatus; **jf**, glossopharyngeal, vagus, and accessory nerves (CN IX, CN X, and CN XI) transiting jugular foramen; **ms**, median sulcus; **ob**, olfactory bulb; **oft**, olfactory tract; **opt**, optic tract (CN II); **otcc**, orbitotemporal canal; **p**, pons; **pfl**, paraflocculus; **pl**, piriform lobe; **rhf**, rhinal fissure; **sof**, oculomotor, trochlear, ophthalmic and maxillary divisions of trigeminal, and abducens nerves (CN III, CN VI, CN V_{1,2}, CN VI) leading to sphenoorbital fissure; **ve**, vermis. Asterisk (*) indicates depression on right lateral piriform lobe.

the foramen ovale, an unusual arrangement unknown in other eutherians. Simpson (1933: fig. 3) argued on the basis of his interpretation of the ventral surface of the endocast of *Te. lucarius* that CN V₂ passed instead through a separate foramen rotundum. The midcranial region of the specimen (AMNH 9245) on which Simpson's (1933) analysis was based shows signs of having been extensively repaired. In any case, neither the skull in its present condition nor the CT scan provides any evidence for an aperture conforming to a foramen rotundum on either side. In addition, the new endocast reconstruction of *Te. lucarius* demonstrates that the casts of CN V₁ and CN V₂ form a single bundle on the left side of the skull, while bone breakage and displacement on the right side gives an artificial impression of two separate bundles (fig. 23C). In short, CN V₂ in *Te. lucarius* must have departed the skull through the sphenoorbital fissure, as in other SANUs.

Foramen ovale, transmitting CN V₃, is large, circular in cross section, and obliquely oriented with respect to the floor of the brainstem (fig. 22A). In *Te. lucarius* (Simpson, 1933) the foramen is more vertically aligned than in IANIGLA-PV 29.

Posterior cranial fossa. The position of the internal acoustic meatus and morphological origins of the facial (CN VII) and vestibulocochlear (CN VIII) nerves are indicated by a small protuberance on the endocast at the level of the anterior portion of the cerebellum_c (fig. 22A). The track of the glossopharyngeal (CN IX), vagus (CN X), and accessory (CN XI) nerves toward the jugular foramen is suggested by a thick, anterolaterally directed feature on the floor of the brainstem. The cast of the canal transmitting the hypoglossal (CN XII) nerve through the single hypoglossal foramen is posterior and slightly medial to the previous structure.

ENDOCRANIAL VASCULATURE: Pneumatization of the skull complicates the identification of some vessels (figs. 10, 11, 17). In dorsal view (fig. 22B), the median sulcus would have supported a large superior sagittal sinus, draining blood from

the dorsal portion of the brain into the transverse sinuses (e.g., Sisson and Grossman, 1975; Hiatt and Gartner, 2001). On the endocast of IANIGLA-PV 29 only a short section of the transverse sinus is preserved as an impression. The transverse sinus connects with (1) the foramina for rami temporales (fig. 17D), indicating vascular communication with the scalp, (2) the posttemporal canal (fig. 17A), transmitting the arteria diploëtica magna and a system of veins related to one or more venous sinuses (Wible, 1987, 2010; MacPhee, 2014), (3) the canal for the postglenoid vein (fig. 17C), via a short sigmoid vessel of very small diameter, and (4) the orbitotemporal canal.

Of these, the track of the orbitotemporal canal is of great interest because it differs from the routing usually seen. On the lateroventral aspect of the endocast, below the piriform lobe_c (fig. 22A, C), there is a very distinct track that must be vascular because it runs along the periphery of the middle cranial fossa, low on the endocranial aspect of the squamosal. This corresponds to the orbitotemporal canal (= sinus canal), which transmits the ramus supraorbitalis (the anterior division of the ramus superior of the stapedial artery) and/or sinus vein (Wible, 1987, 2008). As confirmed by the CT scan, its pathway is partly intracranial and partly intramural, i.e., within the skull's sidewall (fig. 13D, E). On the specimen's left side, the cast of this vessel leaves the endocranial cavity via a small aperture within the orbit, at the level of, but posterior to, the optic foramen (fig. 14). A similar, but more dorsally located, structure is seen on the endocast of *Te. lucarius* AMNH 9245 (Simpson, 1933; fig. 23 A, B). In other regards the track is similar, as it runs forward almost horizontally at the level of the CN II_c. The diameter of the cast of the track is substantial in both specimens.

If our interpretation of conditions in IANIGLA-PV 29 is correct, the orbitotemporal canal in this specimen is unusual, for this feature normally creases the inner sidewall of the skull more dorsally, at the level of the upper portion of the squamosal or even the parietal (e.g., Wible, 1987,

1993; Rougier et al., 1992). This places the vessel against the dorsal half of the piriform lobe (Orliac et al., 2012) or actually on the rhinal fissure (= lateral cerebral venous sinus of Simpson, 1933), for which it is sometimes considered a probable marker (e.g., Rowe, 1996; Silcox et al., 2011; Orihuela, 2014).

UPPER DENTITION

TOOTH MORPHOLOGY

The dental formula is I1–3, C, P1–4, M1–3 as in other macraucheniiids (fig. 24; Bond, 1999). As already noted, the dentition of IANIGLA-PV 29 is exceptionally worn, with teeth in some cases having been worn down to their roots or actually avulsed during life. These factors limit morphological description.

I1–3 alveoli are preserved on each side of the premaxilla and follow the curve of the upper jaw as in *H. cristata* (MLP 41-IV-29-4), *O. zeballosi* (MACN-PV 13671), and *Pr. calchaquiorum* (MACN-PV 5528). The I1s are separated by a median diastema that is wider than the alveoli that it borders. The alveolus for the canine is oval and separated from I3 and P1 by short diastemata, as in *H. cristata* (MLP 41-IV-29-4). The premolars and molars are heavily worn with predominant (and asymmetrical) wear on the left side. Enamel is preserved as a thin layer on the labial and distal surfaces of the better-preserved teeth and along the inner aspect of the fossettes. Of LP1 only its two roots remain, which are similar in size. The P2s are similarly worn, but the posterior root is larger. The RP3, worn down to the base of its crown, is more rectangular in shape in occlusal view than are the anterior premolars. RP4, also quadrangular in outline, exhibits two enamel fossettes on its lingual side, with the mesial the smallest. Both M1s were lost during life and their alveoli have been partially resorbed. The M2s are quadrangular in outline, with two lingual fossettes. The distal fossette is larger and more labially placed than the mesial fossette, similar to conditions in *Pa. denticulata* (MACN-PV 4444). The labial styles are smooth

and the paracone and metacone valleys are moderately pronounced. The M3s are trapezoidal in outline with the mesial portion wider than the distal as in other macraucheniiids. There are three fossettes on the lingual side, in mesial to distal order, with a smaller fossette in a central position (latter only preserved on LM3).

TOOTH HISTOLOGY

The dental histology of litopterns has been discussed for certain Paleogene taxa in relation to enamel microstructure (Line and Bergqvist, 2005). In IANIGLA-PV 29, the dental tissues are well preserved and clearly identifiable on the sectioned M2 (fig. 25). The coronal section, cut at the level of the interroot pad, shows a typical conformation with a succession of enamel (preserved only in a small area labially), dentine, and cementum layers in that order from the occlusal surface to the root (fig. 25A). The distal root transverse section also shows a classical conformation for a senescent non-euhypsodont mammal, showing a nearly obliterated pulp cavity. Covering the root externally, the cementum layer is thin over most of the coronal section, and shows rest lines. However, this layer is notably thickened in the mesial portion of the section, just dorsal to the interroot pad, closer to the crown of the tooth. The existence of this thickened area facilitated the counting and correlation of rest lines in both sections; fig. 25B, D). Growth zones (i.e., between the rest lines) have similar thicknesses, ca. 0.2 mm. Although less clear, the transverse root section also presents a similar number of rest lines. The use of cementum rest lines in age estimation is discussed separately (see Longevity).

DISCUSSION

SYSTEMATICS

TAXONOMY: The hypodigm of *Huayqueriana* presently consists of four specimens. The first is the holotype, MACN-PV 8463 (fig. 2), collected in the Huayquerías de Biluco (De Carles, 1911),

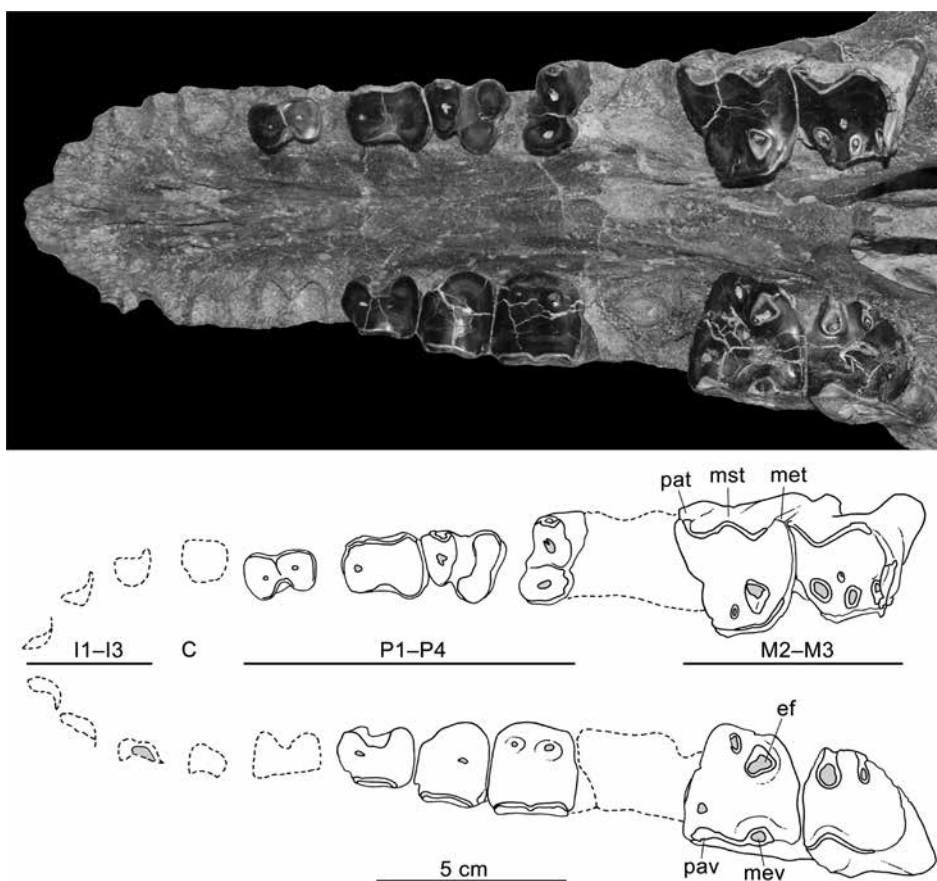


FIG. 24. Palate of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, with detail of dentition in occlusal view. Abbreviations: **C**, canine (alveolus); **ef**, enamel fossette; **I**, incisor (alveolus); **M**, molar; **met**, metastyle; **mev**, metacone valley; **mst**, mesostyle; **P**, premolar; **pat**, parastyle; **pav**, paracone valley.

south of Huayquerías del Este, and believed to have come from the HF (Soria, 1986; Pascual and de la Fuente, 1993). However, as noted earlier (see Geographic and Geological Contexts), the interpretation of the stratigraphic position of the holotype is currently uncertain.

Huayqueriana cristata was originally placed within the genus *Promacrauchenia* (Rovereto, 1914). Kraglievich (1934) later erected the subgenus *Huayqueriana*, which Soria (1986) subsequently elevated to the status of full genus. Soria also considered *Macrauchenidia latidens* (MLP 37-III-7-2), from the Epecuén Formation (Buenos Aires), to be a synonym of *H. cristata* (Soria, 1986). In the same study, Soria (1986) referred

MLP 41-IV-29-4 to *H. cristata* (figs. 5A, 6A, 7A). The fourth and last specimen to be attributed to *Huayqueriana* is IANIGLA-PV 29, which we formally designate as *Huayqueriana* cf. *H. cristata*.

IANIGLA-PV 29 shares some nonexclusive features with the holotype of *H. cristata* (e.g., similar size as recorded in table 2, rostral border of the orbit almost level with the distal border of M3, and convergence of maxillary bones at the level of the P3/P4 embrasure). MACN-PV 8463 (holotype), MLP 41-IV-29-4, and IANIGLA-PV 29 all share possession of a flat snout as seen in lateral view and notably protuberant orbits as seen in dorsal view. IANIGLA-PV 29 also shares with MLP 41-IV-29-4 the round outline of the

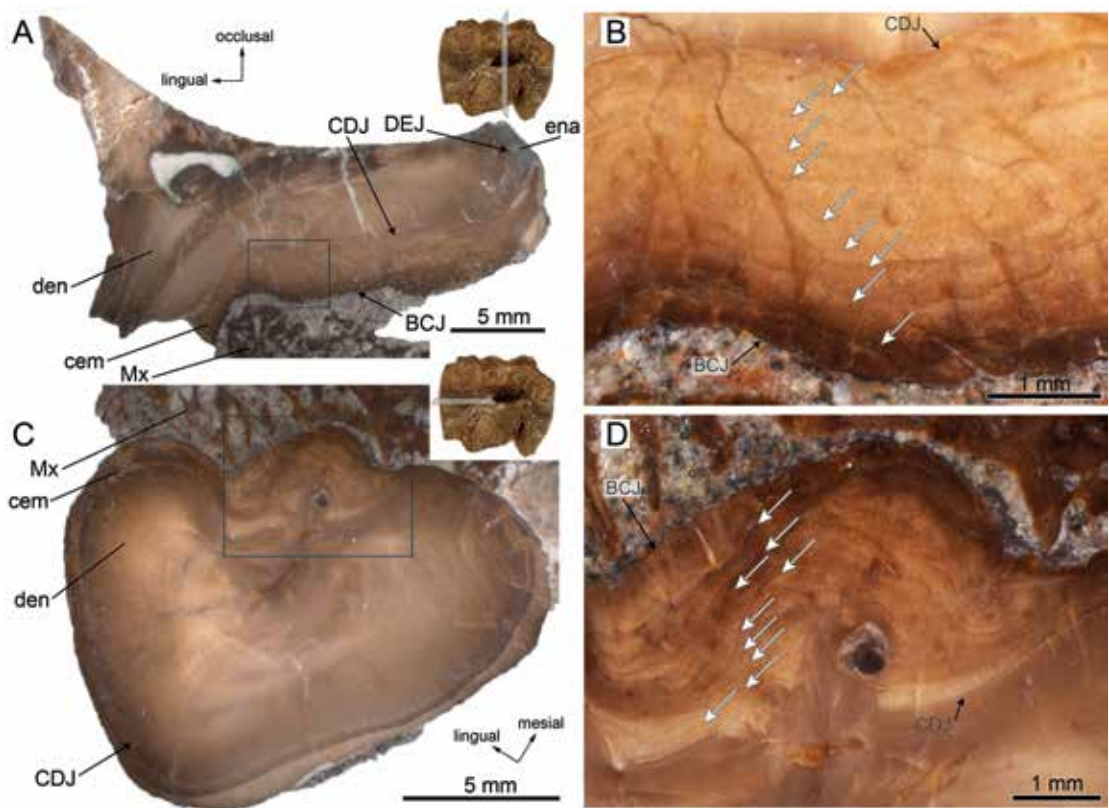


FIG 25. Dental histology of left M2 of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29. Coronal section: **A**, whole surface and **B**, detail of cementum layer (rectangle in **A**). Para-occlusal section: **C**, whole surface and **D**, detail of cementum layer (rectangle in **C**). Cementum rest lines indicated by white arrows. Abbreviations: **BCJ**, bone-cementum junction; **CDJ**, cementum-dentine junction; **cem**, cementum; **DEJ**, dentine-enamel junction; **den**, dentine; **ena**, enamel; **Mx**, maxilla.

premaxillary area in palatal view and small diastemata between I3-C and C-P1. None of these features is identifiable in the holotype.

By contrast, both IANIGLA-PV 29 and MLP 41-IV-29-4 lack the sagittal crest evident in the holotype. To explain this difference, Soria (1986) suggested that MLP 41-IV-29-4 represented an earlier ontogenetic stage than the holotype, arguing that the crest grew with positive allometry during ontogeny.

The sagittal crest provides extra attachment for the temporalis muscle and its production is a direct consequence of the development of this muscle (Holbrook, 2002). In many mammals the growing muscular mass modifies the surface of

the vault such that the temporal lines migrate dorsally and medially to eventually coalesce as the sagittal crest (e.g., *Tapirus pinchaque*; Holbrook, 2002). However, as already noted, IANIGLA-PV 29 is in fact a senile specimen, in which the temporal lines converged but never met.

Sagittal crests are constantly present in some large- and medium-sized herbivores (e.g., *Tapirus terrestris*; Holbrook, 2002), while in others it is a matter of individual variation (e.g., *Tayassu pecari*, *Tragulus javanicus*; AMF personal obs.). The crest on the holotype of *H. cristata* is short and restricted to the posterior vault; IANIGLA-PV 29 is actually very similar, in that the temporal lines are in a nearly identical topographic

position, despite their failure to unite. This suggests that sagittal crest presence in *Huayqueriana* may have varied individually. Sagittal cresting has been best studied in primates (e.g., in New World monkeys, Hershkovitz, 1977). According to notes collected by Napier and Napier (1967) and Osman Hill (1966), cresting is possibly related to age or sex (or both) in chimpanzee, *Pan troglodytes*, and various species of the mona monkey, *Cercopithecus mona*, with older, larger individuals sometimes displaying short crests while in others crests are absent or negligible. Whether such variation occurred in any litoptern is unknown, but it should be considered as a possibility when assessing its value for making taxonomic distinctions.

In conclusion, it is evident that resolving the alpha taxonomy of macraucheniiines will be a major task, due to the number of named forms and their rather uniform skull and dental morphology (e.g., Lydekker, 1894; Soria, 1981). Several macraucheniiine species have been erected on isolated (and thus potentially noncomparable) postcranial elements (e.g., *Ma. ensenadensis*, *Pr. kraglievichi*, *Pr. chapalmalense*, *Pr. (Pseudomacrauchenia) yepesi*, *S. rusconii*; Kraglievich, 1930; Parodi, 1931). Size has often been the most important, if not the only, criterion for distinguishing species (e.g., Ameghino, 1883b; Scott, 1910; Parodi, 1931, Soria, 1981, 1986; Cartelle and Lessa, 1988), which further complicates the picture because the degree of within-group variability is essentially unknown or poorly investigated (see Schmidt, 2013).

However, the significant number of features shared by IANIGLA-PV 29 and the holotype of *Huayqueriana* MACN-PV 8463 and referred specimen MLP 41-IV-29-4 indicates that its referral to this genus is correct. At the same time, there are also some differences that may or may not be due to intraspecific variation. This point can be resolved only by the discovery of additional specimens and further taxonomic studies.

PHYLOGENETIC ANALYSIS: A parsimony analysis was performed based on 19 taxa scored for 34 morphological cranial and dental characters,

using a modified version of the matrix of Schmidt and Ferrero (2014) and eight additional characters (appendix 2). Six characters on which a logical sequence could be imposed to their states were treated as additive (chars. 1, 4, 25–27, 29).

The taxon set includes seven outgroup members: two Adiantidae, including *Proadiantus excavatus* and *Tricoelodus* spp.; and five cram-
aucheniiine macraucheniiids, including *Cram-
auchenia normalis*, *Coniopternium* spp., *Polymorphis lechei*, *Pternoconius* spp., and *Theosodon* spp. The composition of the outgroup varied, depending on the analysis (see below). The ingroup includes 12 macraucheniiine macraucheniiids: *Cullinia levis*, *Huayqueriana cristata*, *Oxydontherium zeballosi*, *Paranauchenia hystata*, *Paranauchenia denticulata*, *Promacrauchenia antiquua*, *Promacrauchenia calchaquorum*, *Scalabrinitherium bravardi*, *Xenorhinotherium bahiense*, *Macrauchenia patachonica*, *Macrauchenopsis ensenadensis*, and *Windhausenian delacroixi*, plus IANIGLA-PV 29 here identified as *Huayqueriana* cf. *H. cristata*.

The data matrix (appendix 2) was analyzed using TNT 1.1 (Goloboff et al., 2008) under maximum parsimony. The analysis included a heuristic search of Wagner trees with 1000 random addition sequences, followed by TBR and retention of 10 trees per round.

The first analysis (all taxa included, *Proadiantus excavatus* used as “strict outgroup” to give plesiomorphic states of characters) yielded 27 most parsimonious topologies (length, 70; CI, 0.643; RI, 0.786; fig. 26A). Macraucheniiinae is supported by 11 characters. The standard bootstrap GC values in our trees are low for all groups (macraucheniiine GC value, 39; fig. 26A), and this analysis failed to satisfactorily resolve the phylogenetic position of macraucheniiines. The consensus tree displays a polytomy for macraucheniiines, with only *Pa. denticulata* and *Pa. hystata* recovered as sister taxa and the Late Pliocene/Pleistocene species *W. delacroixi*, *X. bahiense*, *Ma. ensenadensis*, and *M. patachonica* forming a monophyletic group.

A second analysis was performed using only the taxa for which 50% or more of the characters could be scored, and with *Polymorphis lechei*, *Cramauchenia normalis*, and *Theosodon* spp. as outgroup. *Polymorphis lechei* was used as the “strict outgroup” (fig. 26B). Outgroup taxa incompletely coded were pruned. The analysis provided three trees (length, 61; CI, 0.738; RI, 0.863). Even so, *H. cristata* and *Huayqueriana* cf. *H. cristata* failed to form a monophyletic group, as did the species of *Promacrauchenia*. Standard bootstrap GC values improved for all nodes (macraucheniine GC value, 84; fig. 26B). Forcing the monophyly of *Huayqueriana* and *Promacrauchenia* species required only one additional step.

A third analysis, limited to taxa with 75% or more characters scorable, with only *Cramauchenia normalis* and *Theosodon* spp. as the outgroup (the former used as “strict outgroup”), resulted in three trees (length, 55; CI, 0.818; RI, 0.915; fig. 26C). *Oxydontherium zeballosi* and *S. bravardi* placed respectively as successive sister taxa to the remaining macraucheniines, but the standard bootstrap GC values are very low at their nodes (fig. 26C). This arrangement resembles somewhat the result obtained by Schmidt and Ferrero (2014), with *Scalabrinietherium* as the sister taxon of *Oxydontherium* and remaining macraucheniines. In contrast, Soria (1986) interpreted skull features of *Huayqueriana* as representing the primitive morphology for Macraucheniinae. Our results are inconclusive (fig. 26A, B); however, in the third, most restricted, analysis (fig. 26C), *Huayqueriana* was found to be nested within macraucheniines, closest to *Macrauchenia* and related Plio-Pleistocene taxa. Overall, lack of resolution is probably due either to the choice of characters, or to inherent problems with the group’s alpha taxonomy (e.g., Soria, 1981) that may in turn be largely driven by unappreciated intra-specific variation.

CHARACTER ANALYSIS AND TAXON DEFINITION: Macraucheniinae is a monophyletic group of medium- to large-sized macraucheniids (fig. 26), characterized by the following

combination of features (diagnosis modified from Soria, 1981).

Skull: Long rostrum and maxilla posteriorly and dorsally projecting; reduced nasals (char. 1:2); nasal aperture on top of skull, level with orbit (char. 2:1); palate narrowing at level of P2 or P3 (char. 3:1); postorbital bar with small gap (char. 4:1) or complete; infraorbital foramen anterior to M3 (char. 26:2); contact of premaxilla and maxilla in dorsal view (char. 27:2), with these bones housing a diverticulum (meatus nasi ventralis) not on the direct passageway for air flowing through the nasopharynx; sagittal crest absent or variably present (char. 30:1).

Dentition: Mesodont and selenodont postcanine teeth with labially projecting parastyle on P3–P4 (char. 10:0); deep concavity between labial styles (parastyle, mesostyle, and metastyle) on M1–M3 (char. 11:1); precingulum on M1–M2, slightly lingual or at same level as posterolingual cingulum (char. 14:1); c–p2 obliquely implanted (char. 22:0) and imbricated.

Postcranium: Fused radius/ulna; partly fused tibia and fibula; and tridactyl autopodium.

Most of the character transformations identified on the cranium of macraucheniines are related in one way or another to the massive dorsocaudal retraction and enlargement of the nasal aperture. These transformations, which can be traced in successive taxa over time from Early through Late Miocene (Ameghino, 1893; Scott, 1910, 1937; Rusconi 1957; Soria, 1986), yielded the distinctive “telescoped” skull of later macraucheniines in which the facial bones became caudally displaced relative to the braincase. Among earlier taxa, the stem macraucheniine *Theosodon* (fig. 26) is especially noteworthy for the role it has played in efforts to interpret the highly derived architecture of later forms (Ameghino, 1893; Scott, 1910, 1937; Rusconi 1957; Soria, 1986).

Nasal aperture retraction has invited considerable comment over the years (e.g., Burmeister, 1864, Wall, 1980; Milewski and Dierenfeld, 2013), leading to the hypothesis that it was connected to a mobile proboscis like that of an elephant, a hydrostat like that of a tapir, or, perhaps most inventively, an oper-

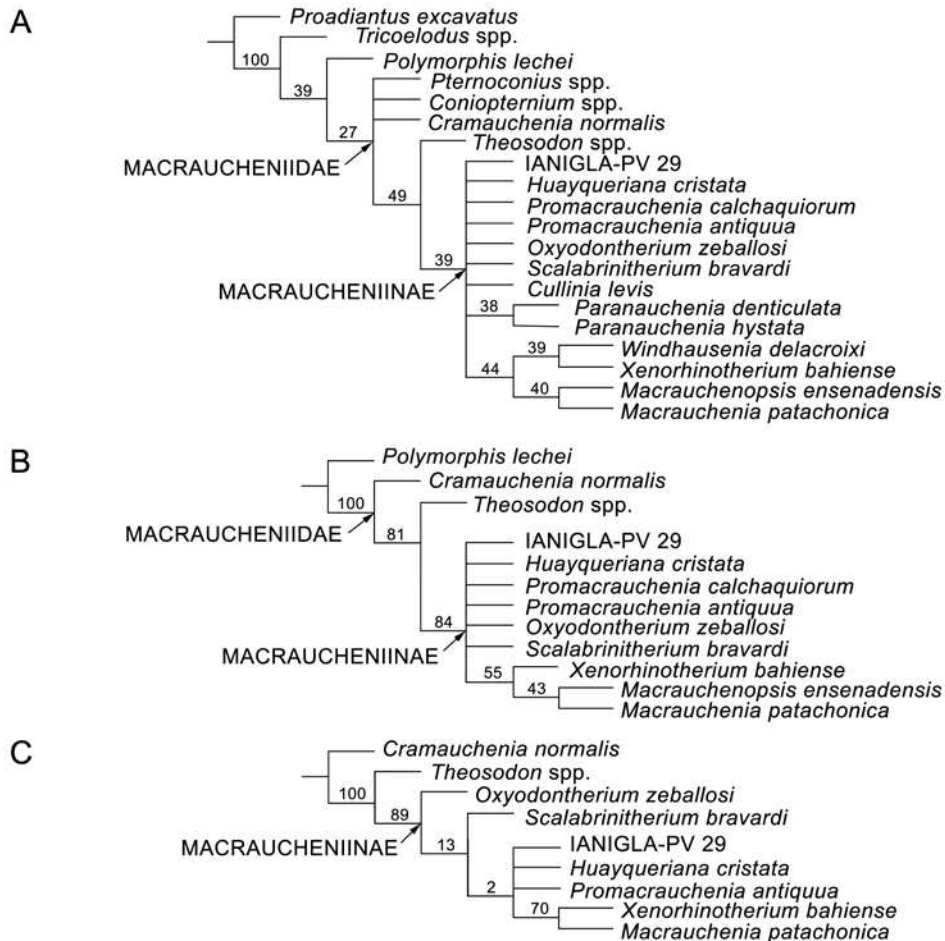


FIG. 26. **A**, Strict consensus of 27 most parsimonious trees (length, 70; CI, 0.643; RI, 0.786); **B**, strict consensus of 3 most parsimonious trees (length, 61; CI, 0.738; RI, 0.863), restricted to taxa with 50% or more of total characters scorable; **C**, strict consensus of 3 most parsimonious trees (length, 55; CI, 0.818; RI, 0.915), restricted to taxa with 75% or more characters scorable. Bootstrap GC values are given at nodes.

culum (Sefve, 1925; Scott, 1937; Rusconi, 1957; Soria, 1981; Bond, 1999). Whatever the nature of associated soft tissues, the progressive reorientation of the air passage in macraucheniines from a position coplanar with the palate to roughly at a right angle to the latter is apparently unique among terrestrial mammals. No other taxa, including elephants (e.g., Todd, 2010), display a comparable level of modification. The only real approach in a morphological sense occurs in odontocetes (e.g., *Tursiops*; Racicot and Colbert, 2002), a morphofunctional convergence that defies any obvious explanation.

PALEOBIOLOGY OF *HUAYQUERIANA*

HEAD POSTURE: Following De Beer (1947), the possible life position of the head of *Huayqueriana* IANIGLA-PV 29 was explored by digitally projecting the 3D model of the inner ear onto the skull, with the plane of the LSC parallel to the horizon (fig. 27). This orientation is biologically meaningful because the LSC is usually horizontal to the substrate when the animal's head is in the position of normal visual alertness (De Beer, 1947; Ekdale, 2013). At rest

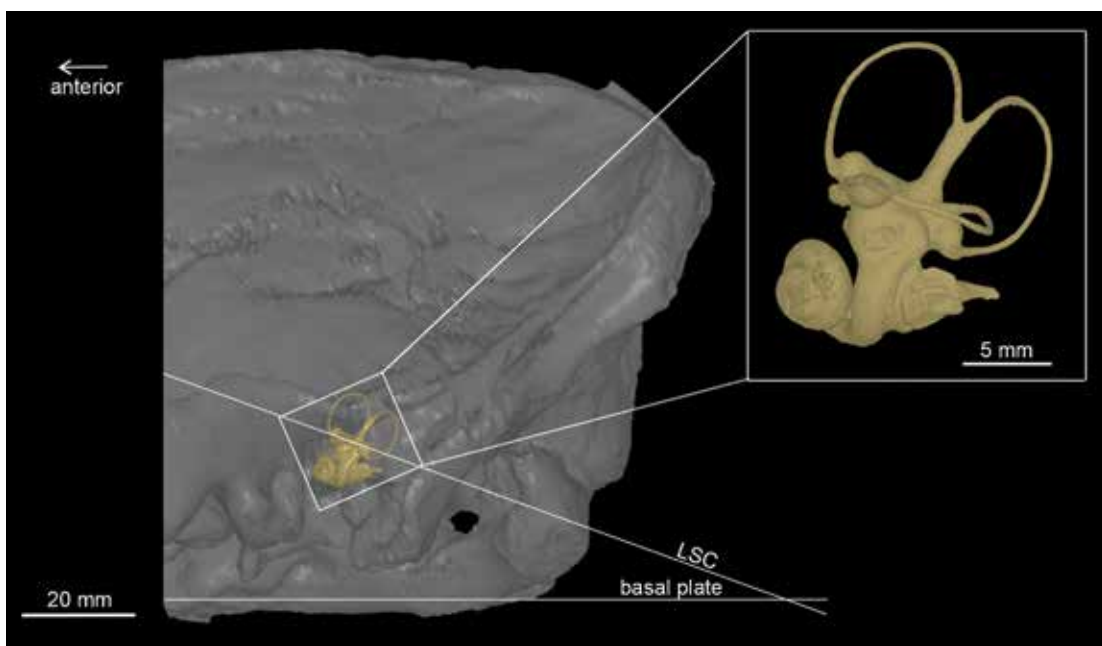


FIG. 27. 3D model of skull and inner ear of *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29, showing angular relationship between plane of lateral semicircular canal (LSC) and that of skull floor (basal plane), following De Beer (1947).

or during dynamic activities (i.e., locomotion, grooming, feeding) the LSC is displaced relative to the horizon (Hullar, 2006). When the LSC is horizontal, the ASC and PSC are vertical and would experience the least stimulation when the head is rotated. This disposition allows the organ of balance to function at maximum efficiency (De Beer 1947).

In IANIGLA-PV 29, the angle subtended by the plane of the LSC and the plane of the base of the skull is about 26° , which is a value similar to that of the horse (De Beer, 1947) (fig. 28). Such a head posture in *Huayqueriana* might have enabled partial binocular vision, as occurs in horses and certain other ungulates (Miller, 1975; Marcus and Sarmiento, 1996). Additionally, among xenarthrans, a strongly inclined LSC was found in armadillos, and regarded as consistent with their “nose down” head position (G. Billet., personal commun., 2016).

BODY SIZE: Our efforts to predict the body mass of IANIGLA-PV 29 using several different

approaches are summarized in table 5. Mean body mass predicted by the equations published by Janis (1990) imply that the mass of *Huayqueriana* cf. *H. cristata* approached 400 kg. This is a significantly larger body mass than known empirical ranges for allometrically comparable panperissodactyls (Welker et al., 2015) such as extant *Tapirus* (*T. indicus*, 250–375 kg; *T. terrestris*, 77–300 kg) and wild *Equus caballus* (200–300 kg) (Nowak and Paradiso, 1983; Silva and Downing, 1995; Clauss et al., 2003). It is also noteworthy that the range of Janis’ estimators is considerable (154 to 721 kg), with a percent prediction error above 25%.

A second approach, utilizing algorithm 4.1 of Mendoza et al. (2006), produces a considerably lower estimate of ~230 kg for IANIGLA-PV 29 (table 5). The percent prediction error was also lower, at 16%. A third approach, using the equations published by Cassini et al. (2012b), yielded the lowest percent prediction error of all (~6%). Their CR4 statistic utilizes the centroid size of a

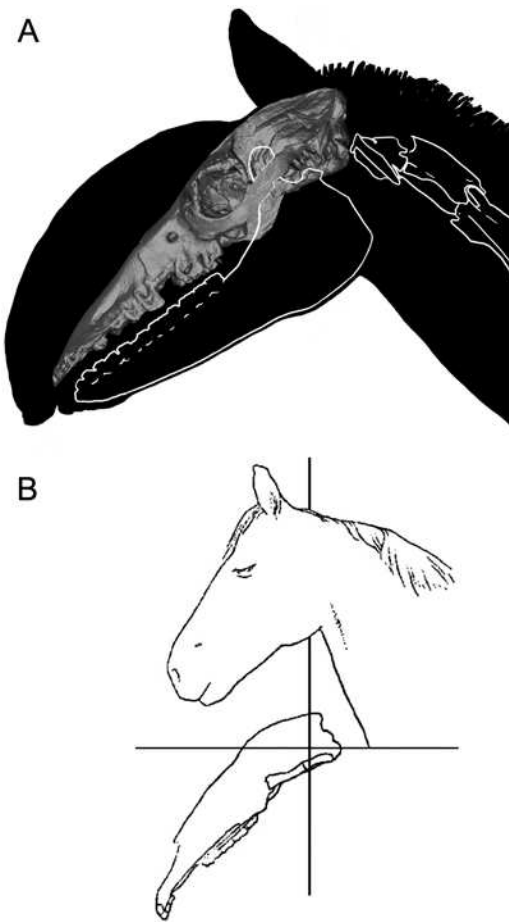


FIG. 28. Reconstruction of orientation of head of **A**, *Huayqueriana* cf. *H. cristata* IANIGLA-PV 29 and **B**, horse (following De Beer, 1947), as they might appear in state of normal visual alertness.

configuration of 36 3D cranial landmarks to predict body mass, which in this case produced a value of 250 kg.

Interpretation of body mass estimates based on different body parts is often difficult because they produce inherently incompatible results. Thus in the case of Janis' (1990) equations, considerable disparity exists between results based on dental vs. cranial material of IANIGLA-PV 29 (table 5), with dentally based estimators yielding substantially larger body sizes. As more comprehensive studies show (Fortelius, 1990; Millien and Bovy, 2010), body

mass estimations based on cranial and postcranial measurements are preferable to ones based on dental measurements.

To assess apparent overestimation in a quantitative manner, Cassini et al. (2012b) developed a quantile regression statistic (Q_{sup}) that provides a theoretical maximum body mass for a given centroid size (CS) based on (in this instance) extant ungulates. In the case of IANIGLA-PV 29, there is no extant ungulate with a CS similar to that of the fossil whose mean body mass surpasses 419 kg (CS maximum, for *Bos taurus*), and in light of this we reject results produced by all three of Janis' dental equations (SUMA, SUMW, SUML). Fariña et al. (1998) reached a similar conclusion when employing Janis' dental regression equations to predict the body size of *Macrauchenia patachonica* (e.g., SUML = 1500 kg vs. their mean estimate of 988 kg based on 66 cranial and postcranial equations). This suggests that a body mass of 400 kg for *Promacrauchenia*, as estimated by Vizcaino et al. (2012), probably also requires reassessment.

By contrast, the mean for Janis' nondental equations is ~265 kg, a value more consistent with results obtained using algorithm 4.1 of Mendoza et al. (2006) and the CR4 equation of Cassini et al. (2012b). In particular, a value of 250 kg for the body mass of *Huayqueriana* IANIGLA-PV 29, using the CR4 statistic of Cassini et al. (2012b), seems more realistic than much larger estimates, and is more in accord with body sizes seen in distantly related taxa such as extant *Tapirus* and *Equus*. These results suggest that a rough doubling of body size occurred in macraucheniniines between the time of Santacrucian *Theosodon* (~130 kg; Cassini et al., 2012a, 2012b) and Huayquerian *Huayqueriana*. Further insights into macraucheniniine body size will require discovery of associated elements of *Huayqueriana*, particularly long bones.

LONGEVITY: Given the extremely advanced dental wear of IANIGLA-PV 29, it can be argued that the specimen was approaching its maximum longevity at the time of death. Counting cementum rest lines, which in principle are deposited

TABLE 5
Body mass estimation of *Huayqueriana* cf. *H. cristata*

Expressions	Prediction error (%)	Body mass (BM) in kg
Janis (1990) ¹ :		
TSL	30.5	339.27
OCH	28.1	154.39
PSL	33.4	261.67
BCL	51.8	306.11
SUML	34.7	407.52
SUMW	38.9	721.18
SUMA	32.7	562.55
Average		393.24
Christiansen and Harris (2005) ² :		
Weighted mean		397.35 ³
Mendoza et al. (2006) ³ :		
Algorithm 4.1	13.5–17.5	228.83
Cassini et al. (2012b) ⁴ :		
CR4		250.6593
Qsup		419.3906

¹ Equations of Janis (1990) after Cassini et al. (2012a): Log10 (BM) = 2.975* log10 (TSL) – 2.344; Log10 (BM) = 2.873*Log10 (OCH) – 0.457; Log10 (BM) = 2.758*Log10 (PSL) – 0.973; Log10 (BM) = 3.137*Log10 (BCL) – 1.062; Log10 (BM) = 3.184*Log10 (SUML) + 1.091; Log10 (BM) = 3.004* Log10 (SUMW) + 1.469; Log10 (BM) = 1.568* Log10 (SUMA) + 1.277; in which BCL, basicranial length; BM, body mass in kg, OCH, occipital height; PSL, posterior skull length; SUMA, second upper molar area; SUML, second upper molar length; SUMW, second upper molar width; TSL, total skull length. Measurements in table 2.

² Equation from Christiansen and Harris (2005) for multiple estimates (*n*). The final body mass (*M*) average is:

$$W_M = \sum_{M_i}^1 ((M_i / \%PE_i) * \sum_{M_i}^1 \%PE) * n^{-2}$$

where *M* is the body mass computed by the regression equations and *n* is the sample size.

³ Equation of Mendoza et al. (2006) after Cassini et al. (2012a) : Ln Algorithm 4.1= 0.736*Ln(SUML) + 0.606*Ln(SUMW) + 0.530*Ln(MZW) + 0.621*Ln(PAW) + 0.741*Ln(SC) – 0.157*Ln(SD) + 0.603; in which MZW, muzzle width; PAW, palatal width; SC= PSL, posterior skull length; SD, depth of the face under the orbit; SUML, second upper molar length; SUMW, second upper molar width. Measurements in table 2.

⁴ Centroid size follows Cassini et al. (2012b) : CR4: Log10 (BM)= 3.165*Log10 (CS) – 6.701; Qsup: Log10 (BM)= 3.321*Log10 (CS) – 6.926.

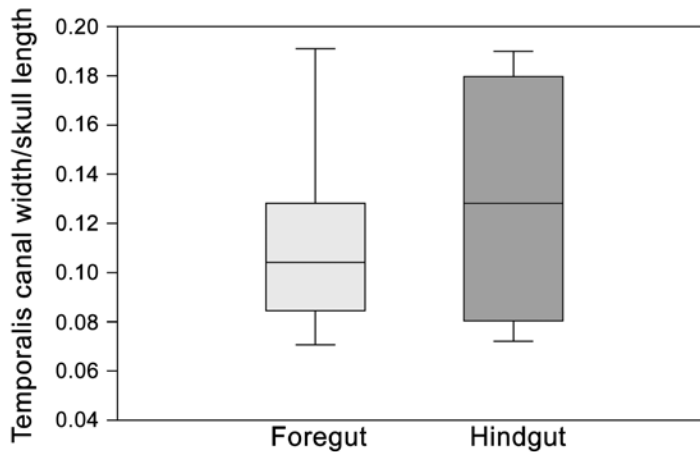


FIG. 29. Box plot depicting ratio of temporalis canal width to skull length in fore- and hindgut fermenters (pooled as such). Foregut and hindgut fermenters are not significantly different, whether data were analyzed all together as here or pooled by family (see fig. 30 and appendix 3).

annually, is often considered the most reliable method of calculating longevity, especially if the animal lives in a temperate climate (Klevezal, 1996) in which there is a distinct low-growth season, usually winter (e.g., Chritz et al., 2009). Age assessment is less precise for mammals living in tropical areas, which lack extreme seasonality. Lines may be indistinct, or two lines may be deposited during a single year (Klevezal, 1996). But given the fact that the Huayquerias fossil site is in a temperate region and that seasonality in the area was probably already established by the Late Miocene, we consider that these lines represent annual depositions. Nine cementum lines can be counted on the sectioned M2 of IANIGLA-PV 29. However, converting that to a plausible age requires correction. Because the last growth layer (external to the last rest line, where the periodontal ligament attached the tooth to the maxilla) is almost as thick as each preceding layer (fig. 25), it may be concluded that the animal died at the end of a year. Based on the data of Klevezal (1996) for the M1 and M2 of ungulates, a correction of 0 to 1 year must be added to the cementum count. Therefore, we suggest that IANIGLA-PV 29 died at an age of almost 10 or 11 years.

Longevity is known to be positively correlated with body size in mammals (Schmidt-Nielsen, 1984). We compared the estimated age of IANIGLA-PV 29 to the longevity of wild extant ungulates approximating a body mass of 250 kg (longevity data from AnAge database, Tacutu et al., 2013; body mass ranges from Nowak and Paradisio, 1983). Longevity is notably greater in the latter. Examples are: ~15 years for the topi, *Damaliscus lunatus*; 24 years for the muskox, *Ovibos moschatus*; and 22 years, or possibly more, for the elk, *Cervus canadensis*, or moose, *Alces alces*. By contrast, ungulates with a longevity in the wild of roughly 10–12 years are conspicuously smaller, with a mean body mass below 100 kg (e.g., the east Caucasian tur, *Capra cylindricornis*, 50 kg; the markhor, *Capra falconeri*, 41 kg; the goitered gazelle, *Gazella subgutturosa*, 49 kg; the Nile lechwe, *Kobus megaceros*, 90 kg; the gerenuk, *Litocranius walleri*, 44 kg; the common rhebok, *Pelea capreolus*, 25 kg; the blue duiker, *Philantomba monticola*, 6 kg; the mountain reedbuck, *Redunca fulvorufula*, 30 kg; the brocket, *Mazama*, 17–33 kg; and the pygmy hog, *Sus salvanius*, 8 kg). The clear conclusion is that, compared to the longevity of modern wild ungulates of a size similar to *Huayqueriana*, the latter had a much shorter lifes-

pan. Note that the magnitude of this conclusion would only increase if larger size estimates were used for *Huayqueriana*.

To provide additional context we also sectioned the M1 of an adult *Theosodon garrettorum* (PIMUZ A/V 4662). Its wear stage is very advanced (styles are completely worn on M1, but on M2–M3 the external outlines are still evident). The specimen exhibited 6 to 7 cementum lines and a thin last growth layer, yielding a likely age of 6–8 years, which is much lower than expected for its body size (ca. 140 kg; Cassini et al., 2012a). Indeed, artiodactyls of similar size live up to ~20 years in the wild (e.g., fallow deer, *Dama dama*, 25 years; Pere David's deer, *Elaphurus davidianus*, 18 years). Our finding opens the question of the influence of environment and food items on longevity reduction in mesodont herbivores during the late Neogene. Unlike notoungulates and many other clades of South American placentals, litopterns failed to develop euhipysodont dentitions as an outcome of adaptation to ecosystem change (Madden, 2015).

DIGESTIVE PHYSIOLOGY: Herbivores utilize different strategies to efficiently digest cellulose-rich vegetation. The broadest partition is between taxa that ferment plant material in the foregut (stomach and/or specialized portions of the small intestine; e.g., ruminant artiodactyls, hippos, colobine monkeys, muroid rodents, sloths, kangaroos, koalas) as opposed to those that utilize the hindgut (caecum and/or colon; e.g., perissodactyls, proboscideans, hyraxes, ateline monkeys, caviomorph rodents, rabbits, wombats) (Clauss et al., 2003; Fletcher et al., 2010). In general, extant hindgut fermenters have a greater capacity for food intake than foregut fermenters of similar size and diet, and also chew their food more on initial ingestion (Clauss et al., 2003; Fletcher et al., 2010). The most specialized foregut fermenters, among which are ruminant artiodactyls, rely on regurgitation, rechewing, and remixing of digestive fluids to ensure maximum extraction of nutrient value from food. In contrast, after initial intake and chewing, hindgut fermenters rely exclusively on chemical reac-

tions in the caecum or colon for nutrient extraction—an ultimately less efficient process.

Although there are marked differences between foregut and hindgut fermenters in digestive physiology, these differences are not reflected osteologically in any obvious way, which makes predictions difficult or impossible for extinct taxa. In principle, differentiation might be expected in the relative occlusal surface area of the cheek teeth. For example, occlusal area seems to be larger in hindgut fermenters of similar size and diet, who also chew food more vigorously in the first stages of digestion than foregut fermenters do (Vizcaíno et al., 2006; Cassini et al., 2012a). Differences in jaw mechanics may also be indicative, at least in theory, as jaw size and musculature should be greater in hindgut fermenters of similar body size and diet than foregut fermenter (e.g., Fletcher et al., 2010).

Based on a study of the occlusal surface areas of cheek teeth, Cassini et al. (2012a) demonstrated that all Early Miocene litopterns may have had a gut physiology comparable to hindgut fermenters. In contrast, Madden (2015) concluded that macraucheniids appear to have possessed an inferred masticatory muscle mass more like that of foregut fermenting ruminants than hindgut digesters. He assessed masticatory muscle mass in 24 specimens (representing 11 taxa of various placental groups) on the basis of two osteological measurements, skull length (SL) and width of temporalis canal or temporal fossa (TS). On his analysis the two fermentation styles were found to form two distinct groups (Madden, 2015).

In order to test Madden's hypothesis statistically, we measured SL and TS in 43 foregut and hindgut fermenters (distributed across 11 taxa; appendix 3). In our analysis, foregut and hindgut fermenters did not form distinct groups (fig. 29). As the data do not follow a normal distribution as judged by the Shapiro-Wilk and Anderson-Darling tests (p -value < 0.01 for the foregut fermenter group), either a nonparametric test or a data transformation is needed to compare the two groups. Using the nonparametric Mann-Whitney U test, we found that there is no signifi-

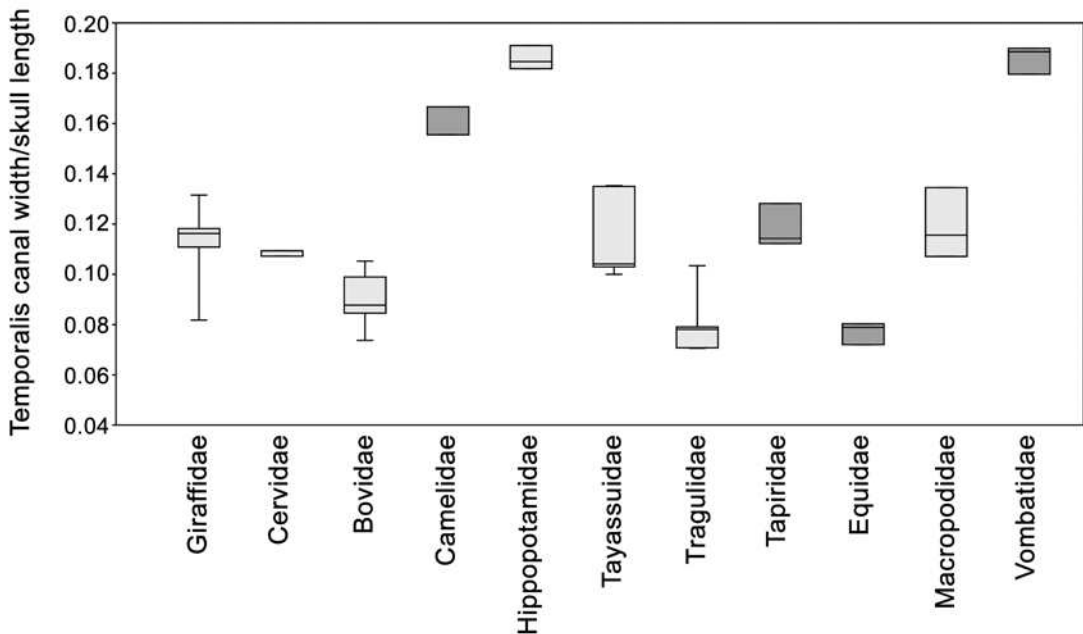


FIG. 30. Box plot depicting ratio of temporalis canal width to skull length, with data pooled by family (appendix 3).

cant difference between the two groups (p -value >0.14). Similarly, after log-transforming the data and checking for both normality of the distribution (Shapiro-Wilk and Anderson-Darling tests, p -value >0.4) and homoscedasticity (Levene's test, p -value >0.10), we found once again that the two groups were not significantly different (Student's t -test, p -value >0.09). We also pooled the data by family in order to compare our results to Madden's analysis (fig. 30), but with the same result. In sum, our results indicate that extant foregut and hindgut fermenters cannot be separated from one another only using skull length and width of the temporal fossa as proxies for masticatory muscle mass.

The hindgut fermentation strategy seems to be the plesiomorphic condition for ungulates (Prothero and Foss, 2007). In view of the fact that there are no foregut fermenters in crown Perissodactyla, the extant group to which litopterns, notoungulates, and possibly other SANUs are most closely related (Welker et al., 2015; Buckley, 2015), phylogenetic bracketing predicts

that litopterns were not either. Further insights may be provided by other predictive models not assessed here (e.g., Cassini et al., 2012a).

CONCLUSIONS

The various families of South American native ungulates grouped as Litopterna thrived on that continent from at least the late Paleogene to the beginning of the Holocene. The terminal taxa of Litopterna included the Macraucheniinae, characterized inter alia by a highly derived cranial structure, in which the nasal aperture was rotated to a position between the orbits, near the summit of the skull. This contribution presents a detailed study of the morphology of a newly discovered and nearly complete macraucheniine skull (IANIGLA-PV 29) from the Huayquerías Formation (Late Miocene, Huayquerian SALMA) in west-central Argentina (Mendoza Province). Referral of IANIGLA-PV 29 to *Huayqueriana* cf. *H. cristata* (Rovereto, 1914) (Litopterna, Macrauche-

niidae, Macraucheniinae) is based on nonexclusive, but overwhelmingly similar, features occurring in *Huayqueriana cristata*.

Phylogenetic analysis failed to satisfactorily resolve the position of IANIGLA-PV 29 within Macraucheniinae, a result possibly influenced by intraspecific variation. Macraucheniinae is a monophyletic group with good support. We have emended the diagnosis of the subfamily to include derived features detected by our phylogenetic analysis.

Thanks to the availability of micro-CT scanning, we have been able to report much new information on the construction of the macraucheniine skull. Chief morphological findings may be grouped under these headings:

1. Reorganization of the nasal cavity: The nasal cavity extends into the rostrum, as in mammals generally, but this rostral portion (meatus nasi ventralis) is now merely a diverticulum of the “true,” vertically oriented air pathway used in life. What this implies for the external soft-tissue structure of the face of *Huayqueriana* and other macraucheniines remains uncertain.

2. Foramina: *Huayqueriana* and its allies lacked a separate foramen rotundum. CT scanning also revealed that the maxillary nerve (CN V₂) leaves the skull through the sphenoorbital fissure, a point previously in contention.

3. Pneumatization: Paranasal airspaces inflate much of the cranial vault, nasal region, and basicranium. By contrast, paratympanic pneumatization is slight.

4. Head orientation: The geometry of the lateral semicircular canal relative to the enclosing skull indicates that the head was habitually held at an appreciable angle, probably much as in modern horses.

5. Internal anatomy: The 3D reconstruction of the endocast, petrosal, and inner ear revealed the presence of several derived features that will provide information for future phylogenetic analyses (e.g., orbitotemporal canal running below the rhinal fissure along the lateroventral aspect of the piriform lobe; stapedia ratio ~1.8;

very shallow subarcuate fossa; coiled cochlea with 2.2 full turns).

6. Body mass: Body mass estimation for fossil taxa, especially those having no close living relatives, presents many difficulties. Although our efforts to establish a plausible size for *Huayqueriana* were inconclusive, we believe that an estimate in the vicinity of 250 kg is much more likely than higher values.

7. Longevity: The extremely advanced dental wear seen in IANIGLA-PV 29 suggests that the animal was at the end of its natural life span. Cementum line counting indicates that it died at 10 to 11 years of age, which is lower than the expected lifespan of extant wild ungulates of similar body mass. This “accelerated aging” may be an outcome of ecosystem change in the Miocene and later, in South America, affecting the longevity of mesodont herbivores.

8. Digestion: Although it has been suggested that some litopterns were probably foregut fermenters (i.e., reliant on regurgitation, rechewing, and remixing of digestive fluids to ensure maximum extraction of nutrient value from food), our evaluation of the method is that it is at best inconclusive. There is no meaningful evidence that any macraucheniines were foregut fermenters.

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APPENDIX 1
COMPARATIVE SET: LITOPTERNA

Family	Species	Specimen	Locality	Stratigraphy	Age/SALMA	Major Reference
Proterotheriidae	<i>Tetramerorhinus lucarius</i> Ameghino, 1894	AMNH 9245	Felton's Estancia; Santa Cruz	Santa Cruz Formation	E. Miocene/Santacrucian	Simpson, 1933 ¹
	<i>Tetramerorhinus cingulatum</i> (Ameghino, 1891b)	MACN-A 5971	La Cueva; Santa Cruz	Santa Cruz Formation	E. Miocene/Santacrucian	Soria, 2001
Macraucheniidae Cramaucheniinae	<i>Cramauchenia normalis</i> Ameghino, 1902	MACN-A 52-219	Gran Barranca; Chubut	Sarmiento Formation	E. Miocene/Colhuehuapian	Dozo and Vera, 2010
	<i>Cramauchenia normalis</i> Ameghino, 1902	MPEF-PV 2524	Cabeza Blanca; Chubut	Sarmiento Formation	L. Oligocene/Deseadan	Dozo and Vera, 2010
	<i>Theosodon lydekkeri</i> Ameghino, 1887b	MACN-A 9269	Corrighuen Aike; Santa Cruz	Santa Cruz Formation	E. Miocene/Santacrucian	Cassini et al., 2012a
	<i>Theosodon garrettorum</i> Scott, 1910	PIMUZ A/V 4662	Locality not indicated, Santa Cruz	Santa Cruz Formation	E. Miocene/Santacrucian	Cassini et al., 2012a
	<i>Macrauchenia patagonica</i> Owen, 1838	MACN-PV 2	Near Salto, Buenos Aires	Pampeano Formation	Pleistocene-Holocene	Burmeister, 1864; Fernández de Álvarez, 1940
Macraucheniidae Macraucheniinae	<i>Huayqueriana cristata</i> (Rovereto, 1914)	MACN-PV 8463 ²	Huayquerías de San Carlos; Mendoza	Huayquerías Formation	L. Miocene/Huayquerian	Soria, 1986
	<i>Huayqueriana cristata</i> (Rovereto, 1914)	MLP 41-IV-29-4	Huayquerías de San Carlos; Mendoza	Huayquerías Formation	L. Miocene/Huayquerian	Soria, 1986
	<i>Huayqueriana cristata</i> (Rovereto, 1914)	MLP 37-III-7-2	Adolfo Alsina; Buenos Aires	Epecuén Formation	L. Miocene/Huayquerian	Soria, 1986
	<i>Macrauchenia patagonica</i> Owen, 1838	MACN-PV 2	Near Salto, Buenos Aires	Pampeano Formation	Pleistocene-Holocene	Burmeister, 1864; Fernández de Álvarez, 1940
	<i>Macrauchenia patagonica</i> Owen, 1838	MLP 12-1424	Arrecifes, Buenos Aires	Pampeano Formation	Pleistocene-Holocene	Sefve, 1925
	<i>Macraucheniopsis ensenadensis</i> (Ameghino, 1888)	MLP 12-1426	Locality not indicated, Buenos Aires	Pampeano Formation	E Pleistocene/Ensenadan	Paula Couto, 1945
	<i>Oxydontherium zeballosi</i> Ameghino, 1883a	MACN-PV 13671	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L. Miocene/"Mesopotamian"	Schmidt, 2013
	<i>Oxydontherium zeballosi</i> Ameghino, 1883a	MACN-PV 17745 ³	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L. Miocene/"Mesopotamian"	Schmidt, 2013
	<i>Oxydontherium zeballosi</i> Ameghino, 1883a	MHIN-UNSL-GEO-V 465a ⁴	Arroyo La Petra; San Luis	Río Quinto Formation	L. Miocene/Huayquerian	Cerdeño et al. 2008

Family	Species	Specimen	Locality	Stratigraphy	Age/SALMA	Major Reference
	<i>Paranauchenia hystata</i> (Cabrera and Kraglievich, 1931)	MLP 29-IX-1-75	Arroyo Chasicó, Buenos Aires	Chasicó Formation	L Miocene/Chasicoan	Schmidt and Ferrero, 2014
	<i>Paranauchenia denticulata</i> (Ameghino, 1891a)	MACN-PV 4444	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L Miocene/“Mesopotamian”	Schmidt, 2013
	<i>Promacrauchenia antiquua</i> (Ameghino, 1887a)	MACN-PV 7986	Monte Hermoso, Buenos Aires	Monte Hermoso Formation	E Pliocene/Montehermosan	Fernández de Álvarez, 1940
	<i>Promacrauchenia calchaquiorum</i> Rovereto, 1914	MACN-PV 5528 ²	Valle de Santa María, Catamarca	“Estratos Araucanos” ⁵	L Miocene-Pliocene/Chasicoan to Montehermosan ⁵	Rovereto, 1914
	<i>Scalabrinitherium bravardi</i> Ameghino, 1883b	MACN-PV 4414 ⁶	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L Miocene/“Mesopotamian”	Schmidt, 2013
	<i>Scalabrinitherium bravardi</i> Ameghino, 1883b	MACN A-1270	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L Miocene/“Mesopotamian”	Schmidt, 2013
	<i>Scalabrinitherium bravardi</i> Ameghino, 1883b	MACN-PV 8903	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L Miocene/“Mesopotamian”	Schmidt, 2013
	<i>Scalabrinitherium bravardi</i> Ameghino, 1883b	MACN-PV 13082	Conglomerado Osífero; Entre Ríos	Ituzaingó Formation	L Miocene/“Mesopotamian”	Rusconi, 1932 ⁷
	<i>Xenorhinotherium bahiense</i> Cartelle and Lessa, 1988	MCL-2644	Gruta dos Ossos Estado da Bahia	N/A	Holocene	Cartelle and Lessa, 1988
	<i>Windhausenien delacroixi</i> Kraglievich, 1930	MACN PV 5301	Uquía, Jujuy	Uquía Formation	L. Plioc / Vorohuean-Sanandresian	Kraglievich, 1930; Reguero et al., 2007

¹ Specimen originally assigned to “*Proterotherium cavum*” Ameghino 1887b, but taxonomy here follows Soria (2001:42).

² Holotype.

³ Young adult, unworn M3.

⁴ Juvenile.

⁵ According to MACN catalog, the stratigraphic location of *Pr. calchaquiorum* is the “Araucana Fm” (an informal unit formed by the Chiquimil, Andalhuala and Corral Quemado formations, ranging from ~12 to ~3 Ma). No precise data is available to determine the age of the specimen (Bonini, com. per. 2015).

⁶ Cast of the holotype.

⁷ We follow Rusconi’s assignment of this partial skull to *S. bravardi*.

APPENDIX 2

PHYLOGENETIC ANALYSIS

Data matrix based on Schmidt and Ferrero (2014), with modifications in wording for certain characters and characters states (characters 1–5, 8, 10–12, 15, 18, 19, 21, 23), plus eight additional characters (characters 26–34).

LIST OF CHARACTERS

ABBREVIATIONS: *, additive character

1. Development of nasals *
 - 0 developed, overhanging the nasal aperture
 - 1 reduced
 - 2 vestigial or absent
2. Nasal aperture, caudal border
 - 0 anterior to orbit
 - 1 level with orbit
3. Anterior palatal shape
 - 0 lateral borders almost parallel
 - 1 narrowing at P2 or P3 level
4. Orbit, postorbital bar *
 - 0 open (large gap)
 - 1 open (small gap)
 - 2 closed
5. Outline of premaxillary area in palatal view
 - 0 acute
 - 1 rounded
6. Premaxillary area in lateral view
 - 0 slightly ventrally curved
 - 1 straight
7. Frontal fossae on caudal margin of nasal aperture
 - 0 absent
 - 1 present
8. Height of tooth crown (HI: crown height/labio-lingual width; Janis, 1988) based on m3 when possible
 - 0 brachyodont (HI<1.45)
 - 1 mesodont (HI= or >1.45)
9. Diastema between I3–C
 - 0 present
 - 1 absent
10. Parastyle on P3–P4
 - 0 labially projected
 - 1 level with mesostyle and metastyle
11. Concavities between lingual styles (parastyle-mesostyle-metastyle) on M1–M3
 - 0 shallow
 - 1 deep
12. Position of hypocone on M1–M2
 - 0 hypocone level with metacone
 - 1 hypocone near protocone, mesial to metacone
13. Position of hypocone vs protocone on M2
 - 0 hypocone labial to protocone
 - 1 hypocone lingual to protocone
14. Anterolingual cingulum (precingulum) on M1–M2
 - 0 noticeably more lingual than posterolingual cingulum
 - 1 slightly lingual or at same level as posterolingual cingulum
15. Hypolophulid on p4
 - 0 shorter than paralophid
 - 1 similar in length to paralophid
16. Entolophid on m1–m2
 - 0 present
 - 1 absent
17. Hypoconulid on m3
 - 0 expanded, forming a third lobe
 - 1 not expanded
18. Entoconid on m1–m2
 - 0 weak (does not reach lingual level of the metaconid)
 - 1 developed (reaches lingual level of metaconid)
 - 2 absent
19. Paralophid on m1–m2
 - 0 level with or lingual to lingual border of metaconid

- 1 labial to lingual border of metaconid
20. Entoconid on m3
0 developed and joined to hypolophid
1 incipient on the hypolophid
2 joined to hypolophid
3 absent
21. Metaconid on m2
0 labial or level with paraconid and entoconid/hypoconulid
1 projects lingual to level of paraconid and entoconid/hypoconulid
22. Implantation of c-p2
0 oblique
1 parallel to mandibular border
23. Trigonid valley on p4
0 larger than talonid valley
1 similar in size to talonid valley
24. Entoconid on p4
0 present
1 absent
25. Length of m2 *
0 small (5–20 mm)
1 medium (21–35 mm)
2 large (36–45 mm)
3 very large (more than 45 mm)
26. Position of coronal plane passing through infraorbital foramen *
0 anterior to M1
1 anterior to M2
2 anterior to M3
3 over M3
27. Snout in dorsal view *
0 premaxillae and maxillae separated sagittally
1 premaxillae sutured sagittally
2 premaxillae and maxillae sutured sagittally
28. Choana, rostral border (= posterior border of palate)
0 level with or anterior to posterior border of M2
1 posterior to M2
29. Orbit, coronal plane passing through rostral border *
0 anterior to posterior border of M3
1 level with posterior border of M3
2 considerably posterior to M3 (equivalent to one tooth-length gap)
30. Sagittal crest
0 present
1 absent
31. Basioccipital, median crest
0 absent or vestigial
1 present
32. Skull shape
0 short and wide
1 long and narrow
33. Occipital condyles
0 protruding at level of nuchal crest in lateral view
1 not protruding at level of nuchal crest in lateral view
34. Odontoid notch
0 U-shaped
1 V-shaped

DATA MATRIX

ABBREVIATIONS: ? missing data; a polymorphic taxa (0 and 1); b polymorphic taxa (2 and 3).

TAXA: The following outgroup taxa includes the scoring of different species of the same genus: *Tricoelodus* spp. (*Tr. bicuspidatus* and *Tr. boliviensis*); *Coniopternium* spp. (*C. andinum* and ?*C. primitivum*); *Pternoconius* spp. (*Pt. polymorphoides* and *Pt. tournoueri*) and *Theosodon* spp. (*Th. lydekkeri*, *Th. lallemanti*, *Th. garrettorum*, *Th. fontanae*, *Th. gracilis*, *Th. patagonicum*, and *Th. karaikensis*).

CHANGES IN SCORING: Character 18, *Xenorhinotherium bahiense*, changed from 1 to 0 (entoconid does not reach the lingual level of the metaconid); character 4, *Huayqueriana cristata*, changed from 2 to ? (uncertain whether postorbital bar was complete; see text); character 9, *Huayqueriana cristata*, changed from 1 to 0 (small diastema in MLP 41-IV-29-4 between I3 and C).

<i>Proadiantus excavatus</i>	??????0??	????100112	0?0a0????	???
<i>Tricoelodus</i> spp.	??????0?1	0011100112	0?010????	???
<i>Polymorphis lechei</i>	??????1	0011000112	01110????	???
<i>Cramauchenia normalis</i>	0000100001	0010101113	0?11000?00	1000
<i>Coniopternium</i> spp.	??????0??	0?10001110	0?010????	???
<i>Pternoconius</i> spp.	??????0??	???001112	01110????	???
<i>Theosodon</i> spp.	1000110001	0100001100	a111111000	1111
<i>Cullinia levis</i>	??????0?0	0?11001101	0011a????	???
<i>Scalabrinitherium bravardi</i>	211?110000	1110001100	0?11122101	1?01
<i>Oxyodontherium zeballosi</i>	211100?010	1110001103	0?11a2200?	0??0
<i>Paranauchenia hystata</i>	?????????	1?1?00?00?	11111????	???
<i>Paranauchenia denticulata</i>	??????0?0	1011?01000	1??12??0?	???
<i>Promacrauchenia antiquua</i>	??11000110	1101101001	001112?101	1101
<i>Promacrauchenia calchaquiorum</i>	211?000?10	1101??????	?????22111	11??
<i>Xenorhinotherium bahiense</i>	2112011111	1101101103	001022??21	?101
<i>Macrauchenia patachonica</i>	2112111101	1011111203	00112b2121	0101
<i>Macrauchenopsis ensenadensis</i>	?11?11?100	1101111203	00113????	???
<i>Windhausenia delacroixi</i>	?1????1???	????10?00?	?0102????1	??01
<i>Huayqueriana cristata</i>	211?100?0?	1111??1???	0???1221aa	?1??
<i>Huayqueriana</i> cf. <i>H. cristata</i>	2111100?0?	11???????	?????22111	0110

APPENDIX 3

SKULL LENGTH AND WIDTH OF TEMPORALIS CANAL IN HINDGUT AND FOREGUT FERMENTERS

Measurements in mm. Abbreviations: SL, length of skull (=condylobasal length); TS, width of temporalis canal (maximum distance between zygomatic arch and adjacent part of the braincase in horizontal plane); *, approximate measurement.

Order	Family	Species	Collection number	Digestive physiology	SL	TS
Artiodactyla	Giraffidae	<i>Giraffa camelopardalis</i>	MMB 12076	Foregut	550	45
Artiodactyla	Giraffidae	<i>Okapia johnstoni</i>	NMB 11009	Foregut	490	50
Artiodactyla	Giraffidae	<i>Okapia johnstoni</i>	NMB 11027	Foregut	480	48
Artiodactyla	Giraffidae	<i>Okapia johnstoni</i>	NMB 10778	Foregut	460	53
Artiodactyla	Giraffidae	<i>Okapia johnstoni</i>	NMB 10917	Foregut	470	50
Artiodactyla	Cervidae	<i>Cervus elephas</i>	ZMUZH 19206	Foregut	373	40
Artiodactyla	Cervidae	<i>Cervus elephas</i>	ZMUZH	Foregut	320	35
Artiodactyla	Bovidae	<i>Saiga tatarica</i>	NMB 10132	Foregut	218	20
Artiodactyla	Bovidae	<i>Saiga tatarica</i>	NMB 10307	Foregut	217	16
Artiodactyla	Bovidae	<i>Saiga tatarica</i>	NMB 10065	Foregut	201	17
Artiodactyla	Bovidae	<i>Saiga tatarica</i>	NMB 1908	Foregut	235	20
Artiodactyla	Bovidae	<i>Bubalus depressicornis</i>	NMB 3268	Foregut	285	25
Artiodactyla	Bovidae	<i>Bubalus depressicornis</i>	NMB 2142	Foregut	293	29
Artiodactyla	Bovidae	<i>Bubalus depressicornis</i>	NMB 3001	Foregut	285	30
Artiodactyla	Camelidae	<i>Camelus bactrianus</i>	NMB 5918	Hindgut	450	70
Artiodactyla	Camelidae	<i>Camelus bactrianus</i>	NMB 5270	Hindgut	480	80
Artiodactyla	Hippopotamidae	<i>Hexaprotodon liberiensis</i>	NMB 10941	Foregut	330	60
Artiodactyla	Hippopotamidae	<i>Hexaprotodon liberiensis</i>	NMB 5297	Foregut	335	64
Artiodactyla	Hippopotamidae	<i>Hippopotamus amphibius</i>	NMB 2767	Foregut	650	120
Artiodactyla	Tayassuidae	<i>Tayassu pecari</i>	NMB 8074	Foregut	250	25
Artiodactyla	Tayassuidae	<i>Tayassu pecari</i>	NMB 8228	Foregut	240	25
Artiodactyla	Tayassuidae	<i>Tayassu pecari</i>	NMB 8418	Foregut	233	24
Artiodactyla	Tayassuidae	<i>Tayassu pecari</i>	NMB 3531	Foregut	207	28
Artiodactyla	Tayassuidae	<i>Tayassu pecari</i>	NMB 3894	Foregut	200	27
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 3809	Foregut	89*	6.3
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 3808	Foregut	85*	6
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 6103	Foregut	89*	6.3
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 3804	Foregut	87*	6.8
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 3795	Foregut	88.5*	7
Artiodactyla	Tragulidae	<i>Tragulus javanicus</i>	NMB 3803	Foregut	87*	9
Perissodactyla	Tapiridae	<i>Tapirus terrestris</i>	NMB 2689	Hindgut	350	40
Perissodactyla	Tapiridae	<i>Tapirus terrestris</i>	NMB 10437	Hindgut	383	43
Perissodactyla	Tapiridae	<i>Tapirus terrestris</i>	NMB 5691	Hindgut	390	50
Perissodactyla	Equidae	<i>Equus grevyi</i>	NMB 5463	Hindgut	560	45
Perissodactyla	Equidae	<i>Equus grevyi</i>	NMB 10873	Hindgut	555	40
Perissodactyla	Equidae	<i>Equus grevyi</i>	NMB 10876	Hindgut	570	45
Marsupialia	Macropodidae	<i>Macropus giganteus</i>	NMB 9437	Foregut	173	20
Marsupialia	Macropodidae	<i>Macropus giganteus</i>	NMB 3609	Foregut	168	18
Marsupialia	Macropodidae	<i>Macropus giganteus</i>	NMB 5453	Foregut	195	25
Marsupialia	Macropodidae	<i>Macropus giganteus</i>	NMB 7993	Foregut	171	23
Marsupialia	Vombatidae	<i>Lasiorhinus latifrons</i>	NMB 10851	Hindgut	167	30
Marsupialia	Vombatidae	<i>Vombatus ursinus</i>	NMB 4148	Hindgut	175	33
Marsupialia	Vombatidae	<i>Vombatus ursinus</i>	NMB 1929	Hindgut	179	34

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